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PCT

NOTIFICATION OF ELECTION

(PCT Rule 61.2)

Assistant Commissioner for Patents United States Patent and Trademark Office

From the INTERNATIONAL BUREAU

Box PCT

Washington, D.C.20231 **ETATS-UNIS D'AMERIQUE**

	217110 01110 2 711112111 202
Date of mailing (day/month/year) . 04 October 2000 (04.10.00)	in its capacity as elected Office
International application No.	Applicant's or agent's file reference
PCT/CA00/00147	DH/12987.5
International filing date (day/month/year)	Priority date (day/month/year)
11 February 2000 (11.02.00)	11 February 1999 (11.02.99)
Applicant	
DESGROSEILLERS, Luc et al	

1.	The designated Office is hereby notified of its election made:
	X in the demand filed with the International Preliminary Examining Authority on:
	06 September 2000 (06.09.00)
	in a notice effecting later election filed with the International Bureau on:
	· · · · · · · · · · · · · · · · · · ·
2.	The election X was
	was not
	made before the expiration of 19 months from the priority date or, where Rule 32 applies, within the time limit under Rule 32.2(b).
	•

The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland

Authorized officer

F. Baechler

Telephone No.: (41-22) 338.83.38

Facsimile No.: (41-22) 740.14.35

PCT



INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

1-

Applicant's of KB/12987	r agent's file refe .5	FOR FURTHER ACTION	See Notification of Transmittal of International Preliminary Examination Report (Form PCT/IPEA/416)
International	application No.	International filing date (day/mo	onth/year) Priority date (day/month/year)
PCT/CA0	0/00147	11/02/2000	11/02/1999
International C12N15/5		ation (IPC) or national classification and IPC	
Applicant UNIVESI	TE DE MONT	REAL et al.	
1. This in and is	iternational pre transmitted to	liminary examination report has been prepa the applicant according to Article 36.	red by this International Preliminary Examining Authority
2. This R	EPORT consis	sts of a total of 9 sheets, including this cove	er sheet.
b€ (s	een amended a ee Rule 70.16	o accompanied by ANNEXES, i.e. sheets of and are the basis for this report and/or sheet and Section 607 of the Administrative Instru- ist of a total of 6 sheets.	f the description, claims and/or drawings which have ts containing rectifications made before this Authority actions under the PCT).
3. This re	eport contains i	ndications relating to the following items:	
1	Basis of the state of	the report	
II.	Priority		
III	☑ Non-esta	ablishment of opinion with regard to novelty,	inventive step and industrial applicability
١٧		unity of invention	
V	□ Reasone citations	ed statement under Article 35(2) with regard and explanations suporting such statement	to novelty, inventive step or industrial applicability;
VI	⊠ Certain o	documents cited	
VII	⊠ Certain d	defects in the international application	
VIII	□ Certain c	observations on the international application	1

Date of completion of this report Date of submission of the demand 04.05.2001 06/09/2000 Authorized officer Name and mailing address of the international preliminary examining authority: European Patent Office - P.B. 5818 Patentlaan 2 Montero Lopez, B NL-2280 HV Rijswijk - Pays Bas Tel. +31 70 340 - 2040 Tx: 31 651 epo nl

Telephone No. +31 70 340 3739



INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No. PCT/CA00/00147

1.	the and	receiving Office in i	nents of the international appresponse to an invitation under this report since they do no	ler Article 14 are	referred to in this	report as "originally filed"
	1-32	2	as originally filed			
	Clai	ms, No.:				
	1-38	3	as received on	13/04/2001	with letter of	11/04/2001
	Dra	wings, sheets:				
	1/20)-20/20	as originally filed			
2.	With lang	n regard to the lang Juage in which the	juage , all the elements mark international application was	ed above were a filed, unless othe	vailable or furnish erwise indicated u	ed to this Authority in the nder this item.
	The	se elements were a	available or furnished to this	Authority in the f	ollowing language	: , which is:
		the language of a	translation furnished for the	purposes of the i	nternational searc	h (under Rule 23.1(b)).
		the language of pu	ublication of the international	application (und	er Rule 48.3(b)).	
		the language of a 55.2 and/or 55.3).	translation furnished for the	purposes of inter	national prelimina	ry examination (under Rule
3.			eleotide and/or amino acid ry examination was carried o			
		contained in the in	ternational application in wri	tten form.		
		filed together with	the international application	in computer read	dable form.	
		furnished subsequ	ently to this Authority in writ	ten form.		
		furnished subsequ	iently to this Authority in com	nputer readable f	orm.	
			t the subsequently furnished pplication as filed has been t		e listing does not	go beyond the disclosure in
		The statement that listing has been full	t the information recorded in irnished.	computer reada	ble form is identic	al to the written sequence
4.	The	amendments have	e resulted in the cancellation	of:		
		the description,	pages:			
		the claims,	Nos.:			



International application No. PCT/CA00/00147

		the drawings,	sheets:
5.	Ø		established as if (some of) the amendments had not been made, since they have been ond the disclosure as filed (Rule 70.2(c)):
		(Any replacement sh report.) see separate sheet	eet containing such amendments must be referred to under item 1 and annexed to this
6.	Add	litional observations, i	necessary:
III.	Non	n-establishment of o	pinion with regard to novelty, inventive step and industrial applicability
1.			e claimed invention appears to be novel, to involve an inventive step (to be non- ally applicable have not been examined in respect of:
		the entire internation	al application.
	×	claims Nos. 29.	
be	caus	se:	
			application, or the said claims Nos. relate to the following subject matter which does ational preliminary examination (<i>specify</i>):
			ns or drawings (indicate particular elements below) or said claims Nos. are so unclear pinion could be formed (specify):
	⊠	the claims, or said cl opinion could be form	aims Nos. 29 are so inadequately supported by the description that no meaningful ned.
		no international sear	ch report has been established for the said claims Nos
2.	and	eaningful internationa /or amino acid sequer ructions:	Il preliminary examination cannot be carried out due to the failure of the nucleotide nce listing to comply with the standard provided for in Annex C of the Administrative
			not been furnished or does not comply with the standard.
		the computer readab	le form has not been furnished or does not comply with the standard.
IV.	Lac	ck of unity of invention	on
1.	In re	esponse to the invitati	on to restrict or pay additional fees the applicant has:
		restricted the claims.	





International application No. PCT/CA00/00147

	\boxtimes	paid additional fees.										
		paid additional fees und	er prote	st.								
		neither restricted nor pa	id additi	onal fees								
2.		This Authority found tha 68.1, not to invite the ap						nplied and	l chose,	accordi	ing to Ru	ule
3.	This	s Authority considers that	the req	uirement	of unity o	of invention i	n accorda	ance with	Rules 1	3.1, 13.2	2 and 13	3.3 is
		complied with.										
	×	not complied with for the see separate sheet	e followii	ng reasor	ıs:							
4.		nsequently, the following (mination in establishing t			ational a	application w	ere the s	ubject of i	nternatio	onal prel	liminary	
	☒	all parts.										
		the parts relating to clair	ns Nos.									
V.		asoned statement under tions and explanations					, inventi	ve step o	r indust	rial app	olicabilit	ty;
1.	Stat	tement										
	Nov	velty (N)	Yes: No:	Claims Claims	2, 5-7							
	Inve	entive step (IS)	Yes: No:	Claims Claims	2, 5-7							
	Indu	ustrial applicability (IA)	Yes: No:	Claims Claims	2, 5-7							

VI. Certain documents cited

2. Citations and explanations see separate sheet

1. Certain published documents (Rule 70.10)

and / or

2. Non-written disclosures (Rule 70.9)

see separate sheet



INTERNATIONAL PRELIMINARY EXAMINATION REPORT



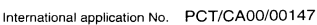
International application No. PCT/CA00/00147

VII. Certain defects in the international application

The following defects in the form or contents of the international application have been noted: see separate sheet

VIII. Certain observations on the international application

The following observations on the clarity of the claims, description, and drawings or on the question whether the claims are fully supported by the description, are made: see separate sheet





INTERNATIONAL PRELIMINARY Inte

Re Item I Basis of the report

The amendments filed with the International Bureau under Article 19(1) introduce subject-matter which extends beyond the content of the application as filed, contrary to Article 19(2) PCT. The amendments concerned are the following:

- 1. Claim 1: The application as filed does not disclose a method for obtaining a Neprilysin-like metallopeptidase as claimed in claim 1. The application discloses in pages 14-16 a particular method for the cloning and recombinant expression of mouse NL1 of which claim 1 constitutes an undue generalization.
- 2. Claims 3, 4, and 5-7 as far as dependent on claims 3 and 4: A soluble metallopeptidase sharing about 80% homology with the C-terminus of the furin site shown in Figure 3 has not been specifically disclosed in the application. Soluble forms of NL1 and NL3, consisting of the ectodomain are referred to in page 10. This does not provide support for soluble enzymes sharing 80% homology with the disclosed soluble forms.
- 3. Claims 8 and 22: The application as filed does not disclose such a general method for obtaining a substrate of a metallopeptidase. Page 24 discloses several methods for identifying substrates, of which claims 8 and 22 constitute an undue generalization.
- 4. Claims 9-12 and 23-26: Page 25 discloses a method for identifying inhibitors. The method of claims 9 and 23 constitutes an undue generalization of the subject-matter disclosed in page 25. No support exists in the application for inhibitors or uses thereof other than the speculative statement referred to in page 25.
- 5. Claims 13 and 27: The direct of use of the metallopeptidase to manage disease is not disclosed in the application. Page 27 discloses merely that it is possible to help the patient by managing the activity of the enzyme.
- 6. Claims 14, 15 and 28: The application discloses a sequence encoding the N-terminal part up to the furin-recognition sequence in figure 3 (see page 10), but not a sequence

having 80% homology to it.

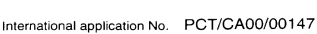
- 7. Claims 16-21: A metallopeptidase sharing about 80% homology with the C-terminus of the furin site shown in Figure 4 has not been specifically disclosed in the application. Soluble forms of NL1 and NL3, consisting of the ectodomain are referred to in page 10. This does not provide support for soluble enzymes sharing 80% homology with the disclosed soluble forms.
- 8. Claims 30-38: A metallopeptidase sharing about 80% homology with the C-terminus of the transmembrane domain shown in Figure 5 has not been specifically disclosed in the application. Soluble forms of NL1 and NL3, consisting of the ectodomain are referred to in page 10. This does not provide support for soluble enzymes sharing 80% homology with the disclosed soluble forms, neither to any of its applications.

Re Item IV

Lack of unity of invention

The present application relates to endopeptidase-like metallopeptidases and a method for obtaining. Polypeptides belonging to the neutral metallopeptidase family have been already been described in the state of the art (see pages 1 and 2 of the description). The article "Gene" 1996, vol.174, pages 135-143 discloses the cloning of a metallopeptidase of the neutral endopeptidase family. In the light of the prior art a problem underlying the present application can be formulated as providing further endopeptidase-like metallopeptidases. The following solutions are proposed:

- 1. A polypeptide of the sequence disclosed in figure 3, fragments and variants thereof and corresponding nucleic acid (Claims 2-15).
- 2. A polypeptide of the sequence disclosed in figure 4, fragments and variants thereof and corresponding nucleic acid (Claims 16-27).
- 3. A polypeptide of the sequence disclosed in figure 5, fragments and variants thereof and corresponding nucleic acid (Claims 30-38).



EXAMINATION REPORT - SEPARATE SHEET

A further problem identified in the application relates to a method for obtaining a Neprilysin-like metallopeptidase. The solution proposed as formulated in claim 1 is as follows:

1. Using primers in the C- and N-terminus of the sequence His-Glu-Xaa-Xaa-His sequence in a PCR method.

Given the essential difference between the problems posed and their corresponding solutions, since neutral endopeptidase-like metallopeptidases have already been disclosed in the state of the art, and due to the differences in primary sequence between the polypeptides disclosed as solutions to the first problem, as well as among their corresponding nucleic acid sequences and since in the light of the state of the art, no other technical feature could be distinguished as being new and common to the identified problems and corresponding solutions, the IPEA is of the opinion that there is no single inventive concept underlying the plurality of the claimed inventions in the present application, in the sense of Rule 13.1 PCT.

Re Item V

Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

Reference is made to the following document:

D1: US-A-5 688 640 (MASASHI YANAGISAWA) 18 November 1997 (1997-11-18)

- 1. Claims 2 and claims 5-7, as far as dependent on claim 2 relate to a metallopeptidase sharing about 80% homology with the amino acid sequence of figure 3. No such sequence has been disclosed in the state of the art and therefore, claims 2 and 5-7 are novel and comply with the requirements of Article 33(2) PCT.
- 2. Document D1 discloses the aminoacid and nucleic acid sequences of the Endothelin Converting Enzyme ECE-1 which is a membrane-bound neutral metalloprotease

EXAMINATION REPORT - SEPARATE SHEET

expressed in endothelial cells, with a sequence significantly different from the one disclosed in figure 3 of the application. No hint exists in the state of the art which would allow the skilled person to retrieve a metallopeptidase with 80% homology to the one disclosed in figure 3. Consequently, the subject-matter of claims 2 and 5-7 involves an inventive step and meets the requirements of Article 33(3) PCT.

Re Item VI

Certain documents cited

Certain published documents (Rule 70.10)

Application No Patent No Publication date

(day/month/year)

Filing date (day/month/year)

Priority date (valid claim) (day/month/year)

PCT/FR99/00807

21/10/1999

07/04/1999

08/04/1998

Re Item VII

Certain defects in the international application

- 1. Contrary to the requirements of Rule 5.1(a)(ii) PCT, the relevant background art disclosed in the document D1 is not mentioned in the description, nor is this document identified therein.
- 2. Claim 2 contains a reference to the drawings. According to Rule 6.2(a) PCT, claims should not contain such references except where absolutely necessary, which is not the case here.

Re Item VIII

Certain observations on the international application

1. The term "about" used in claim 2 is vague and unclear and leaves the reader in doubt as to the meaning of the technical features to which it refers, thereby rendering the definition of the subject-matter of said claim unclear (Article 6 PCT).

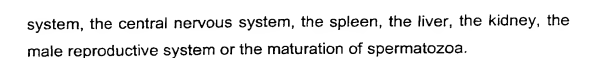
13. 04. 2001

WHAT IS CLAIMED IS:

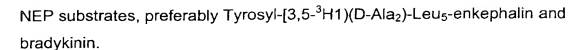
(82)

- A method for obtaining a Neprilysin-like (NEP-like) metallopeptidase which comprises the following steps:
- selecting a primer in C-terminus of the His-Glu-Xaa-Xaa-His (where Xaa represents any amino acid) with a degenerate nucleotide sequence complementary to at least the Gly-Glu-Asn-Ile-Ala-Asp amino acid sequence of known NEP-like metallopeptidases with sufficient binding capacity;
- selecting a primer in N-terminus of the His-Glu-Xaa-Xaa-His (where
 Xaa represents any amino acid) with a degenerate nucleotide sequence complementary to a conserved amino acid sequence with preferably 80% homology with known NEP-like metallopeptidases and sufficient binding capacity;
 - contacting said primer with tissue nucleic acids to yield PCR products;
- selecting said PCR products that contain the His-Glu-Xaa-Xaa-His motif; and
 - completing the sequence of said selected PCR products with standard methods.
- 2. A metallopeptidase sharing about 80% homology with the amino acid sequence shown in Figure 3.
 - 3. A metallopeptidase which is soluble sharing about 80% homology with the amino acid sequence in C-terminus of the furin site shown in Figure 3.
- 4. A metallopeptidase which is soluble sharing about 80% homology with the amino acid sequence shown in Figure 3 and with an enzymatic activity capable of degradation of known Neprilysin substrates, preferably Tyrosyl-[3,5-3H1)(D-Ala₂)-Leu₅-enkephalin and bradykinin.
 - 5. A composition comprising a metallopeptidase as defined in any one of claims 2-4.

- 6. A nucleic acid encoding a metallopeptidase as defined in any one of claims 2-4.
- 7. An antibody directed against a metallopeptidase as defined in any one of claims 2-4.
- 5 8. A method for obtaining a substrate of a metallopeptidase as defined in any one of claims 2-4, which comprises the steps of:
 - -- contacting said metallopeptidase with a molecule or extract; and
 - -- assaying the resulting solution for a decrease in said molecule or extract, when compared with the same but in absence of said metallopeptidase, as an indication of the presence of said substrate.
 - 9. A method for obtaining an inhibitor of a metallopeptidase as defined in any one of claims 2-4, which comprises the steps of:
 - -- contacting said metallopeptidase with a molecule or extract in the presence of a substrate selected known NEP substrates, preferably Tyrosyl-[3,5-3H1)(D-Ala₂)-Leu₅-enkephalin and bradykinin; and
 - -- assaying the resulting solution for an increase in said substrate, when compared with the same but in absence of said metallopeptidase, as an indication of the presence of said inhibitor.
 - 10. An inhibitor obtained from the method of claim 9.
- 20 11. The use of a known NEP inhibitor or an inhibitor as defined in claim 10 to control the enzymatic activity of a metallopeptidase as defined in any one of claims 2-4.
- 12. The use of a known NEP inhibitor or an inhibitor as defined in claim 10 to manage disease relating to the physiological status of the cardiovascular system, the central nervous system, the spleen, the liver, the kidney, the male reproductive system or the maturation of spermatozoa.
 - 13. The use of a metallopeptidase as defined in any one of claims 2-4 to manage disease relating to the physiological status of the cardiovascular



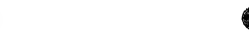
- 14. A recombinant vector comprising 80% homology with the nucleic acid encoding the N-terminal part of the amino acid sequence shown in Figure 3, which N-terminal part terminates with a furin-recognition sequence.
- 15. A method for producing a soluble form of a protein, polypeptide or part thereof which comprises:
 - obtaining nucleic acids encoding said protein, polypeptide or part thereof;
- fusing said nucleic acids in phase with an N-terminal fragment wherein said N-terminal fragment comprises a cleavable furin-like site located in C-terminus past the transmembrane region or is an N-terminal part as defined in claim 14:
 - having the fused nucleic acids to be expressed in a host cell which expresses or is made to express furin in the presence of a culture medium; and
 - recovering said soluble form in the culture medium.
- A protein, polypeptide or part thereof produced by the method defined in claim 15, wherein said protein, polypeptide or part thereof is a metallopeptidase sharing about 80% homology with the region in C-terminus of the putative furin site of the amino acid sequence shown in Figure 4.
- 17. A metallopeptidase sharing about 80% homology with the region in Cterminus of the putative furin site of the amino acid sequence shown in 25 Figure 4.
 - 18. A metallopeptidase sharing about 80% homology with the region in C-terminus of the putative furin site of the amino acid sequence shown in Figure 4 and with an enzymatic activity capable of degradation of known



- 19. A composition comprising a metallopeptidase as defined in any one of claims 16-18.
- 5 20. A nucleic acid encoding a metallopeptidase as defined in any one of claims 16-18.
 - 21. An antibody directed against a metallopeptidase as defined in any one of claims 16-18.
- 22. A method for obtaining a substrate of a metallopeptidase as defined in any one of claims 16-18, which comprises the steps of:
 - -- contacting said metallopeptidase with a molecule or extract; and
 - assaying the resulting solution for a decrease in said molecule or extract, when compared with the same but in absence of said metallopeptidase, as an indication of the presence of said substrate.
- 15 23. A method for obtaining an inhibitor of a metallopeptidase as defined in any one of claims 16-18, which comprises the steps of:
 - -- contacting said metallopeptidase with a molecule or extract in the presence of a substrate selected from known NEP substrates or a protein, polypeptide or part thereof produced by the method of claim 15, preferably Tyrosyl-[3,5-3H1)(D-Ala₂)-Leu₅-enkephalin and bradykinin; and
 - -- assaying the resulting solution for an increase in said substrate, when compared with the same but in absence of said metallopeptidase, as an indication of the presence of said inhibitor.
 - 24. An inhibitor obtained from the method of claim 23.
- 25 25. The use of a known NEP inhibitor or an inhibitor as defined in claim 24 to control the enzymatic activity of a metallopeptidase as defined in any one of claims 16-18.



- 26. The use of a known NEP inhibitor or an inhibitor as defined in claim 24 to manage disease relating to the physiological status of the cardiovascular system, the central nervous system, the spleen, the liver, the kidney, the male reproductive system or the maturation of spermatozoa.
- The use of a metallopeptidase as defined in any one claims 16-18 to manage disease relating to the physiological status of the cardiovascular system, the central nervous system, the spleen, the liver, the kidney, the male reproductive system or the maturation of spermatozoa.
- 28. A method as defined in claim 15, wherein said protein, polypeptide or part thereof is beta-endorphin.
 - 29. A recombinant host cell capable of expressing a protein, polypeptide or part thereof transplanted in a mammal to manage a disease, physiological process or pain.
- 30. A metallopeptidase sharing about 80% homology with the amino acid sequence located in the C-terminus of the predicted transmembrane domain of the amino acid sequence shown in Figure 5 which has been produced by the method of claim 15, by fusing in frame a cleavable signal peptide in N-terminus of said amino acid sequence or by transforming said predicted transmembrane domain into a cleavable signal peptide.
- 20 31. A composition comprising a metallopeptidase as defined in claim 30.
 - 32. An antibody directed against a metallopeptidase as defined in claim 30.
 - 33. A method for obtaining a substrate of a metallopeptidase as defined in claim 30, which metallopeptidase shares about 80% homology with the C-terminal region of the predicted transmembrane domain of the amino acid sequence shown in Figure 5, comprising the steps of:
 - -- contacting said metallopeptidase with a molecule or extract; and



- -- assaying the resulting solution for a decrease in said molecule or extract, when compared with the same but in absence of said metallopeptidase, as an indication of the presence of said substrate.
- 34. A method for obtaining an inhibitor of a metallopeptidase sharing about 80% homology with the C-terminal region of the predicted transmembrane domain of the amino acid sequence shown in Figure 5, which comprises the steps of:
 - -- contacting said metallopeptidase with a molecule or extract in the presence of a substrate produced by the method of claim 33; and
- -- assaying the resulting solution for an increase in said substrate, when compared with the same but in absence of said metallopeptidase, as an indication of the presence of said inhibitor.
 - 35. An inhibitor obtained by the method of claim 34.
- The use of an inhibitor as defined in claim 35 to control the enzymatic activity of the metallopeptidase sharing about 80% homology with the Cterminal region of the predicted transmembrane domain of the amino acid sequence shown in Figure 5.
- The use of an inhibitor as defined in claim 35 to manage disease relating to the physiological status of the central nervous system, the spleen or the bones.
 - 38. The use of a metallopeptidase as defined in claim 30 to manage disease relating to the physiological status of the cardiovascular system, the central nervous system, the spleen or the bones.



(PCT Article 18 and Rules 43 and 44)

Applicant's or agent's file reference DH/12987.5	FOR FURTHER see Notification of (Form PCT/ISA/2)	f Transmittal of International Search Report (20) as well as, where applicable, item 5 below.
International application No.	International filing date (day/month/year)	(Earliest) Priority Date (day/month/year)
PCT/CA 00/00147	11/02/2000	11/02/1999
Applicant		
UNIVESITE DE MONTREAL et	al	- And the same of
This International Search Report has been according to Article 18. A copy is being tra	n prepared by this International Searching Auth ansmitted to the International Bureau.	ority and is transmitted to the applicant
	of a total of sheets. a copy of each prior art document cited in this	report.
	international search was carried out on the bas ess otherwise indicated under this item.	is of the international application in the
the international search w Authority (Rule 23.1(b)).	as carried out on the basis of a translation of th	e international application furnished to this
was carried out on the basis of the contained in the internation	e sequence listing: nal application in written form.	ternational application, the international search
	rnational application in computer readable form this Authority in written form.	1.
	this Authority in computer readble form.	
the statement that the sub international application a	sequently furnished written sequence listing do s filed has been furnished.	pes not go beyond the disclosure in the
the statement that the info furnished	ormation recorded in computer readable form is	identical to the written sequence listing has been
	nd unsearchable (See Box I).	
3. X Unity of invention is lack	king (see Box II).	
4. With regard to the title,		
the text is approved as su	bmitted by the applicant.	
X the text has been establish METALLOPROTEASES OF TH	hed by this Authority to read as follows: IE NEPRILYSIN FAMILY	
- Mari		
 With regard to the abstract, the text is approved as sul 	bmitted by the applicant	
the text has been establish	thed, according to Rule 38.2(b), by this Authority date of mailing of this international search repo	
6. The figure of the drawings to be publi	shed with the abstract is Figure No.	10
as suggested by the applic		None of the figures.
because the applicant faile	ed to suggest a figure. characterizes the invention.	
Decause this ligure better	Characterizes the invention.	



hternational application No. PCT/CA 00/00147

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
see additional sheet
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1, 5-8, 11-22 partially, and 2

Neutral endopeptidase-like metallopeptidase of figure 3, nucleic acid encoding it, vector and host cell comprising the same and use thereof for producing the metallopeptidase; oligonucleotides and antibodies thereof as well as their use for detecting the metallopeptidase in a sample; vector comprising a sequence encoding the N-terminal part of the metallopeptidase and use thereof for producing a soluble form of a protein of interest; soluble metallopeptidase

2. Claims: 1, 5-8, 11-22 partially, 3

Neutral endopeptidase-like metallopeptidase of figure 4, nucleic acid encoding it, vector and host cell comprising the same and use thereof for producing the metallopeptidase; oligonucleotides and antibodies thereof as well as their use for detecting the metallopeptidase in a sample; vector comprising a sequence encoding the N-terminal part of the metallopeptidase and use thereof for producing a soluble form of a protein of interest; soluble metallopeptidase

3. Claims: 1, 5-8, 11, 12, 17-22 partially, 4

Neutral endopeptidase-like metallopeptidase of figure 5, nucleic acid encoding it, vector and host cell comprising the same and use thereof for producing the metallopeptidase; oligonucleotides and antibodies thereof as well as their use for detecting the metallopeptidase in a sample; soluble metallopeptidase

4. Claims: 9, 10

Method for screening molecules related to neutral endopeptidase by using consensus sequences on either side of a His-Glu-Xaa-Xaa-His sequence

PCT/CA 00/00147

Box III TEXT OF THE ABSTRACT (Continuation of item 5 of the first sheet)

Line 2 of the text:

Please delete the word "new".

ernational Application No CT/CA 00/00147

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C12N15/57 C12N9/64

C12N15/62

C07K16/40

C12N15/85 G01N33/573 C12N5/10

C12Q1/68

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B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

C12N C12Q C07K G01N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, WPI Data, BIOSIS, CHEM ABS Data, MEDLINE, STRAND, EMBL

Category °	Citation of document, with indication, where appropriate, of t	the relevant passages	Relevant to claim No.
À	US 5 688 640 A (MASASHI YANAGI 18 November 1997 (1997-11-18) column 2, line 65 -column 6, column 11, line 4 -column 14, column 14, line 64 -column 15 column 17, line 8 -column 18, examples 1-4	line 21 line 35 , line 14	1-8, 11-22
'A" docum consi 'E" earlier	ther documents are listed in the continuation of box C. ategories of cited documents: tent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international date	"T" later document published after the in or priority date and not in conflict wit cited to understand the principle or t invention "X" document of particular relevance; the cannot be considered novel or cannot be co	temational filing date in the application but heory underlying the claimed invention
filing date L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) O* document referring to an oral disclosure, use, exhibition or other means P* document published prior to the international filing date but later than the priority date claimed	"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is taken alon cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family		
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	ll September 2000		

ernational Application No

C.(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	SCOTT I C ET AL: "Molecular cloning, expression and chromosomal localization of a human gene encoding a 33 kDa putative metallopeptidase (PRSM1)" GENE: AN INTERNATIONAL JOURNAL ON GENES AND GENOMES, GB, ELSEVIER SCIENCE PUBLISHERS, BARKING, vol. 174, no. 1, 26 September 1996 (1996-09-26), pages 135-143, XP004043253 ISSN: 0378-1119 the whole document	1-8, 11-22
X	NIGEL M. HOOPER: "Families of Zinc metalloproteases" FEBS LETTERS, vol. 354, no. 1, 31 October 1994 (1994-10-31), pages 1-6, XP002147067 AMSTERDAM NL cited in the application page 1, right-hand column, paragraph 2 page 1, right-hand column, last paragraph -page 2, left-hand column, paragraph 1 page 5, right-hand column, paragraph 1	9,10
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P,X	KOJI IKEDA ET AL.: "Molecular identification and characterization of novel membrane-bound metalloprotease, the soluble secreted form of which hydrolyzes a variety of vasoactive peptides" JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 274, no. 45, 5 November 1999 (1999-11-05), pages 32469-32477, XP002140284 MD US cited in the application abstract page 32469, left-hand column, paragraph 1 -right-hand column, paragraph 1 page 32470, left-hand column, paragraph 3 page 32470, right-hand column, paragraph 1 page 32470, right-hand column, paragraph 4 -right-hand column, paragraph 2	1,5-9, 17-20,22

rnational Application No

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C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT				
ategory °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.		
Э, Х	VALDENAIRE O ET AL: "XCE, A NEW MEMBER OF THE ENDOTHELIN-CONVERTING ENZYME AND NEUTRAL ENDOPEPTIDASE FAMILY, IS PREFERENTIALLY EXPRESSED IN THE CNS" MOLECULAR BRAIN RESEARCH, NL, ELSEVIER SCIENCE BV, AMSTERDAM, vol. 64, no. 2, 1999, pages 211-221-221, XP000863086 ISSN: 0169-328X abstract page 211, left-hand column, paragraph 1 -page 212, right-hand column, paragraph 1 page 212, right-hand column, last paragraph -page 213, left-hand column, paragraph 1 page 215, left-hand column, paragraph 2 -page 216, right-hand column, paragraph 1; figure 1	1,4-9, 19,20,22		
(page 219, left-hand column, paragraph 2 page 219, right-hand column, paragraph 2 -page 220, left-hand column, paragraph 1 & DATABASE EMBL 'Online! Accession number Y16187, 7 January 1999 (1999-01-07) VALDENAIRE, O.: "Homo sapiens mRNA for XCE protein" the whole document	1,4-8, 17-22		

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ernational Application No CT/CA 00/00147

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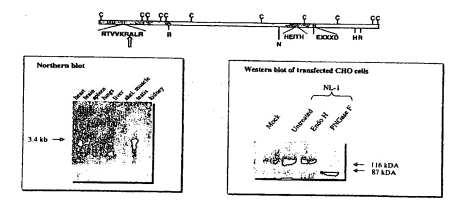
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(54) Title: NEW METALLOPROTEASES OF THE NEPRILYSIN FAMILY

Structure and expression of NL-1



(57) Abstract

In this paper, we describe RT-PCR strategies that allowed us to identify and clone members of the NEP-like family. Degenerate oligoncleotide primers corresponding to consensus sequences located on either side of the HEXXH consensus sequence for zincins were designed and used in RT-PCR with mouse and human testis cDNAs. DNA fragments with lengths expected from the sequence of this class of enzympes were obtained. These DNA fragments were cloned and sequenced. Using this PCR strategy and the PCR fragments as probes to screen cDNA libraries, three zincin-like peptidases were identified in addition of known members of the family. The cDNA sequences allowed to derive specific probes for Northern and in situ hybridization, and probe human chromosomes to localize the gene and establish potential links to genetic diseases. Furthermore, these cDNA sequences were used to produce recombinant fusion proteins in Escherichia coli in order to raise specific antibodies. Finally, the cDNA sequences were cloned in mammalian expression vectors and transfected in various mammalian cell lines to produce active recombinant enzymes suitable for testing specific inhibitors.

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TITLE OF THE INVENTION

New Metalloproteases of the Neprilysin Family

BACKGROUND OF THE INVENTION

Peptides are used by cells from yeast to mammals to elicit physiological responses. The use of peptides as messengers usually involves the following steps: 1) production and release of the peptide by a specific cell, 2) interaction of the peptide with a receptor on the surface of the target cell, and 3) degradation of the peptide to terminate its action. The first and last steps of this scheme require the participation of proteases/peptidases. There is increasing evidence that membrane-associated zincmetallopeptidases play important roles in both of these steps. Although activation of prohormone precursors into bioactive peptides is generally performed by proteases of the subtilisin family located in the Trans-Golgi Network or in secretory granules of the cell (for a review see: (Seidah and Chrétien, 1995)) a few peptides need a final processing step. This step involves the action of membrane-associated zincmetallopeptidases. Two cases are particularly well documented: angiotensinconverting enzyme (ACE) which cleaves inactive angiotensin I into angiotensin II (Corvol and Williams, 1997) and endothelin-converting enzymes (ECEs) which cleave isoforms of big endothelins into endothelins (Turner, 1997a). In addition to their role in peptide activation, cell surface zinc-metallopeptidases have also been implicated in the termination of the peptidergic signal by breaking down the active peptides into inactive fragments. One of the best known of these peptidases is probably Neutral Endopeptidase-24.11 (Neprilysin, NEP) that has been implicated in the physiological degradation of several bioactive peptides (Kenny, 1993). Interestingly, NEP and the ECEs show significant structural similarities and appear to be members of a family of peptidases that also includes PEX, a newly discovered and not yet characterized peptidase, and the KELL blood group protein (Turner and Tanzawa, 1997b). Because of their important role as regulators of bioactive peptide activity, these enzymes (more specifically NEP and the ECEs) have been identified as putative targets for therapeutic intervention, similar to the way ACE inhibitors are used to control blood pressure. The recent discovery of PEX, another member of the family, which appears to be involved in phosphate homeostasis, raised the possibility that other yet unknown members might exist.

Members of the NEP-like family are type II membrane proteins consisting of three distinct domains: a short NH2-terminal cytosolic sequence, a single transmembrane region, and a large extracellular or ectodomain responsible for the catalytic activity of the enzyme. There are potential N-glycosylation sites and cysteine residues that are involved in disulfide bridges stabilizing the conformation of the active enzyme. These enzymes are metalloenzymes with a Zn atom in their active site. As

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such, they belong to the zincin family of peptidases which is characterized by the active site consensus sequence HEXXH (Hooper, 1994), where the two histidine residues are zinc ligands. In members of the NEP-like family of peptidases, the third zinc ligand is a glutamic acid residue located on the carboxy-terminus side of the consensus sequence. This characteristic puts them in the gluzincin sub-family (Hooper, 1994). The model enzyme for gluzincins is thermolysin (TLN) a bacterial protease whose 3D structure has been determined by X-ray crystallography (Holmes and Matthews, 1982). The active site of NEP has been extensively studied by site-directed mutagenesis and several residues involved in zinc binding (Devault et al., 1988b; Le Moual et al., 1991; Le Moual et al., 1994), catalysis (Devault et al., 1988a; Dion et al., 1993), or substrate binding (Vijayaraghavan et al., 1990; Beaumont et al., 1991; Dion et al., 1995; Marie-Claire et al., 1997) have been identified (for a recent review see Crine et al., 1997).

SUMMARY OF THE INVENTION

Here, we developed an RT-PCR strategy to look for other members of this important family of peptidases. This strategy allowed the molecular cloning and characterization of three additional NEP-like (NL) metallopeptidases (called NL-1, NL-2 and NL-3). Knowledge obtained through these studies allows the generation of reagents (nucleic acid probes and primers, antibodies and active recombinant enzymes) for further biochemical characterization of these enzymes and their pattern of expression and will greatly help the rational design of specific inhibitors that could be used as therapeutic agents.

Accordingly, the present invention relates to the following products:

- 25 A. Degenerate primers for screening new NEP-related enzymes;
 - B. NL-1, NL-2 and NL-3 proteins as NEP-related enzymes;
 - C. Nucleic acids encoding these enzymes;
 - D. Antibodies directed against the enzymes;
- E. Recombinant vectors comprising the nucleic acids encoding the enzymes and hosts transformed therewith:
 - F. Fragments of the nucleic acids useful as probes or primers to hybridize and detect the presence of an NL-1, NL-2 and NL-3 genes, or to hybridize and amplify and produce gene fragments;
 - G. Soluble forms of NL-1, NL-2 and NL-3; and
- H. Nucleic acids comprising the N-terminal part of NL-1 or NL-2 which terminates with a sequence encoding a furin recognition site, such nucleic acids being useful for making a fusion protein with the ectodomain of any protein of interest, and for releasing a soluble form of that protein of interest (containing the ectodomain) in the medium.

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Also the present invention relates to the following methods:

- A. A method for screening NEP-related enzymes that make use of degenerate primers or probes selected from a region of NEP family members in a highly conserved region, namely around the zinc-binding sites; and
- 5 B. A method for producing NL-1, NL-2 or NL-3 that includes the steps of culturing the above recombinant host and recovering NL-1, NL-2 and NL-3 gene products therefrom.

The present invention will be described hereinbelow by referring to specific embodiments and appended figures, which purpose is to illustrate the invention rather than to limit its scope.

In the first section, general procedures leading to the identification and localization of NL-1, NL-2 and NL-3 are given. In the second section, slightly different procedures are given for completing or reiterating the work performed on NL-1.

BRIEF DESCRIPTION OF THE FIGURES

Figure 1: Amino acid sequence comparison of human NEP, PEX, KELL and ECE1 peptidases. Amino acid sequences in boxes are those used to design the oligonucleotide primers. Numbers and arrows under the sequences identify the primer and its orientation.

- Figure 2: Sequences of the oligonucleotide primers used in the PCR reactions.
- **Figure 3:** Nucleotide and amino acid sequence of the mouse NL-1 cDNA. The sequence of the DNA fragment obtained by PCR is in brackets.
- **Figure 4:** Partial nucleotide and amino acid sequence of the human NL-2 cDNA. The sequence of the DNA fragment obtained by PCR is in brackets.
- Figure 5: Partial nucleotide and amino acid sequence of the human NL-3 cDNA.
- Figure 6: Amino acid sequence comparison of NEP, NL-1, NL-2 and NL-3 peptidases.
 - Figure 7: In situ hybridization of mouse testis sections using NL-1 as a probe.
 - Figure 8: In situ hybridization of mouse sections using mouse NL-3 as a probe.
 - Figure 9: In situ hybridization of mouse spinal chord sections
 - Figure 10: Expression of NL-1 in mammalian cells.
 - Figure 11: Activity of recombinant soluble NL-1.
- Figure 12: Expression of a soluble form of NL-3 using NL-1 amino-terminal domain.

DESCRIPTION OF THE PREFERRED EMBODIMENTS OF THE INVENTION

SECTION 1)

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MATERIALS AND METHODS

DNA and RNA manipulations

All DNA manipulations and Northern blot analysis were performed according to standard protocols (Ausubel et al., 1988; Sambrook et al., 1989).

<u>mRNA purification and cDNA synthesis</u>

mRNAs were prepared from mouse testis using Quick Prep Micro mRNA purification kit (Pharmacia Biotech). Purified mRNAs were kept at -70° until ready used. First strand cDNA was synthesized from 1µg of mRNA using the First-Strand cDNA synthesis kit (Pharmacia Biotech). The human testis cDNA library was obtained from Clonetech.

Polymerase chain reaction protocol

PCR was performed in a DNA thermal cycler with 5 μ I of cDNA template and 1 μ I of Taq DNA polymerase in a final volume of 100 μ I, containing 1 mM MgCl₂, 2 μ M of each primer oligonucleotide, 20 μ M of each dNTP and 5% DMSO. Cycling profiles included an initial denaturation step of 5 min at 94°C, followed by 30 cycles of 1 min at 94°C, 1 min at 40°C and 1.5 min at 72°C. A final extension step was performed at 72°C for 10 min. The amplified DNA was loaded on a 2% agarose gel and visualized by staining with ethidium bromide. Fragments ranging in size between 500-700 bp were cut and eluted from the gel. If needed, a second round of PCR was done with nested oligonucleotide primers, using 10 μ I of the first PCR reaction, or of the eluted band cut from the agarose gel. Resulting fragments were ligated in pCR2.1 vector (Invitrogen) according to the distributor's recommendations. DH5 α Escherichia coli cells were transformed with the ligation mixture and grown on 2YT plates in the presence of kanamycin. Plasmids were prepared from resistant cells and sequenced. In situ hybridization on mouse tissues and chromosomal localization of human genes

In situ hybridization on whole mouse slices or isolated tissues was performed as described previously (Ruchon et al., 1998).

To determine the chromosomal localization of human NL-2 and NL-3 genes, a technique for mapping genes directly to banded human chromosomes was used. Metaphase chromosomes were obtained from lymphocytes cultured from normal human peripheral blood. Cells were synchronized with thymidine and treated with 5-bromodeoxyuridine (BrdU) during the last part of the S phase to produce R-banding. Biotin-labeling of the probe was done by nick-translation (Bionick, BRL) and the probe was visualized by indirect immunofluorescence. Antibody production

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To raise antibodies against the new peptidases, the cDNA sequences of each protein was compared to that of other members of the family and the sequence segment showing the less homology was used. These sequences are from amino acid residues 273 to 354 for NL-1, from 75 to 209 for NL-2 and from 143 to 465 for NL-3. These cDNA fragments were cloned in vector pGEX2T (Pharmacia Biotechnology) downstream from and in phase with Gluthatione-S-transferase (GST). Plasmids were transformed in E. coli strain AP401 and, induction of synthesis and purification of the fusion proteins were performed as recommended by the supplier. The NL polypeptides were cleaved from the fusion protein with thrombin and purified by SDS-PAGE. NL polypeptides were injected to rabbits or mice according to the following schedules: for rabbits, initial injection of 150 µg of protein with boosts of the same amount 4 weeks and 8 weeks following the initial injection; for mice, initial injection of 100 µg of protein followed by boosts of the same amounts 3 and 6 weeks later. A month after the last injection, sera were collected from the animals and tested by immunoblotting against the initial E. coli-produced antigens and the recombinant proteins produced in mammalian cell lines.

Production of monoclonal antibodies

cDNA fragments corresponding to amino acids segments of NLs selected to raise antibodies were used to construct a GST-fusion protein in E. coli. This fusion protein was purified from E. coli extracts by affinity chromatography on a glutathione-Sepharose column according to the supplier's instructions (Amersham-Pharmacia). After thrombin cleavage, the NL portion of the GST fusion protein was further purified by electroelution from a polyacrylamide gel. This material was used to immunise 4 mice (5 injections of ≈50 µg of NL polypeptide). Blood was collected from each mice after the immunisation schedule and the presence of antibodies in mice serum was assessed by ELISA using microtiter plates coated with NL polypeptide from E. coli extracts. Mice sera were also tested for the presence of NL antibodies by Western blotting extracts of mammalian cells transfected with the NL expression vectors. One mouse selected for its high titer of NL specific antibodies (as measured by ELISA) was sacrificed and its spleen cells were collected and immortalised by fusion with myeloma cells(strain: P3-X63Ag.653 from ATCC) as described previously (Crine 1985). Hybridoma cells were selected for their ability to grow in HAT selection medium and cloned by several rounds of limiting dilution. Hybridomas showing proper affinity and specificity to the enzymes NL-1, NL-2 and NL-3 where selected.

Expression of NLs in cultured mammalian cells and enzymatic assays

The cDNAs for NL-1 and NL-3 were cloned in vectors pcDNA3 or pRcCMV (Invitrogen) and introduced by transfection in mammalian cell lines according to procedures already described in our laboratory (Devault et al., 1988a). Procedures to

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prepare extracts of cellular proteins or culture media were also described in previous papers (Devault *et al.*, 1988a; Lemay *et al.*, 1989). The presence of NLs in these extracts was monitored by immunoblotting using specific antibodies.

Extracts of cellular proteins and culture media were assayed for enzymatic activity. Two tests were performed. The first used [³H]-Tyr-(D)Ala₂-Leu-enkephalin as substrate and was performed according to Lemay et al., (1989). The second used bradykinin as substrate and was performed as described by Raut et al. (1999).

RESULTS

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Cloning of NL-1, a new member of the NEP family

The molecular cloning in the past few years of ECEs, PEX and KELL showed that all these proteins have between 50 and 60% similarity with NEP. This observation led us to believe that these peptidases are part of an extended family and that there could be still additional members to be discovered. To test this hypothesis, we aligned the amino acid sequences of the members of the NEP-like family and designed degenerate oligonucleotide primers to be used in RT-PCR reactions (Figure 1 and 2). These primers were located on either side of the HEXXH consensus sequence for zincins. Because they are highly degenerate, primers 1 and 2 were each subdivided into two pools, 1A-1B, and 2A-2B, respectively (Figure 2). Any PCR amplified DNA fragment that corresponds to a peptidase of the family should normally contain the consensus sequence and be easily recognized by sequencing of the cloned fragments. Using this strategy, we first performed PCR reactions with primer pairs 1A-3 and 1B-3. The amplified DNA migrates mostly as a smear starting at around 700 bp and going down to 100 bp. As the expected fragments should be around 550 bp, we isolated from the gel the section corresponding to DNA fragments longer than 500 bp. A second round of PCR reactions was performed with both crude PCR products of the first reaction and isolated DNA bands, using primers 2A-3 and 2B-3. The expected 296 bp fragment was seen on the gel (not shown).

Cloning of these DNA fragments generated approximately 350 clones, of which 44 were sequenced. Nine of these had no inserts or corresponded to sequences not related to the NEP family, 24 corresponded to NEP, 3 to PEX, and 8 corresponded to one putative new member of the family, since they all contained the HEXXH consensus sequence for zincins and showed 65% homology with mouse NEP (in boxes Figure 3). This fragment was then used to screen a mouse testis cDNA library, and allowed us to isolate a complete cDNA of 2592 nucleotides (Figure 3). The identity of this sequence with other members of the family is presented in Table I. This new member was called NL-1, for NEP-like peptidase 1.

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Cloning of NL-2 and NL-3.

A strategy similar to that described for amplification of enzymes of the NEP family from mouse testis cDNAs was used with a human testis cDNA library using two different oligonucleotide primers. This time, DNA fragments of approximately 900 bp were obtained and cloned. Ten clones were sequenced, revealing the presence of NEP and two new peptidases of the family that we have called NL-2 and NL-3.

The NL-2 PCR fragment was 879 nucleotides in length and encoded a 293 amino acid residue segment probably located in the carboxy-terminal domain of this putative peptidase (in brackets Figure 4). This PCR fragment was then used to screen a lambda gt10 human brain cDNA library. It allowed the isolation of other cDNA fragments which overlap partially with the NL-2 PCR fragment. Fusion of these lambda clones and the PCR fragment resulted in an open reading frame of 770 amino acid residues. The use of 5' RACE protocols with human testis cDNA libraries allowed completion of the sequence of NL-2 ORF (Figure 4). This ORF codes for a putative protein that is about 80% identical to the mouse NL-1 protein (Figure 6). Across species, members of the NEP, PEX, ECEs sub-families have highly conserved sequences (more than 94% identity). Although a sequence identity of about 80% only exists between the novel human protein and mouse NL-1, these proteins share unique characteristics that make possible the fact that NL-2 protein may be the human homologue of NL-1. The identity of NL-2 with other members of the family is presented in Table I.

The 879 bp PCR fragment encoding NL-3 showed an open reading frame of 293 amino acid residues (Figure 5, in brackets). Sequence analysis of NL-3 showed that it was 94.2 % identical to an EST sequence from mouse embryonic tissue present in publicly accessible DNA data banks. This mouse EST sequence, commercially available from American Tissue and Cell Culture (ATCC), had been obtained previously by our laboratories.

Since Northern blot analysis of human tissues with the NL-3 PCR fragment showed the expression of this protein in spinal chord (see below), the same PCR DNA fragment was used to screen by hybridization a human spinal chord cDNA library constructed in phage λ vectors. One clone contained a full-length ORF of 752 amino acid residues that encompassed the 293 amino acid residue ORF of the PCR fragment. Further probing, cloning and sequencing lead to the obtention of NL-3 full sequence, shown in Figure 5.

Figure 6 presents a comparison of the amino acid sequence of the new NEP-like enzymes and Table I shows the extent of identity between members of the family. Cellular distribution of NL-1, NL-2 and NL-3 peptidases

Determining the tissue distribution of NL-1, NL-2 and NL-3 may provide clues to identify the peptidergic systems in which they are involved. It will be particularly

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interesting to compare the tissue distribution of these peptidases with that of NEP and the ECEs to determine whether or not the physiological functions of NL-1 and/or NL-2 and/or NL-3 may overlap those of NEP and/or ECEs.

In situ hybridization (ISH), using our mouse cDNA, was used to determine the spatial and temporal expression of NL-1 during mouse development, as done previously for PEX (Ruchon et al., 1998)). Serial sections of whole foetal (12, 15 and 19 dpc) and adult mice (1, 3 and 6 days old) were hybridized with an [35S]-labeled RNA probe. Figure 7 shows a section of mouse testis which was the only tissue identified to express NL-1 by this technique. Cells of seminiferous tubules are specifically labeled but spermatids located near the center of the tubule showed strongest labeling. These cells are in the last stage of maturation into spermatozoids. The presence of NL-1 in testis has now been confirmed by Northern analysis of mouse tissues (see Fig. 10). Other tissues express NL-1, when analyzed by RT-PCT, which is a more sensitive assay (not shown).

A similar approach was used to determine the localization of NL-3 using the mouse EST obtained from ATCC. Figure 8 shows sections of whole mouse at 17 days of embryonic development and 4 days post-natal. Several tissues are expressing this putative peptidase including brain, where it is associated with neurons (Figure 9), spinal chord, liver, spleen and bones. Labeling was stronger in bones from *Hyp* mouse, an animal model for hypophosphatemic rickets (Figure 8). In bones, NL-3 was found to be expressed by osteoblasts (not shown).

Northern blotting experiments were performed on several tissues with NL-2 and NL-3 probes. A Human Multiple Tissues Northern Blot (Clontech) was hybridized with specific probes. A single RNA band of approximately 4.0 kb was revealed by the probe for NL-2. Expression of NL-2 is restricted to brain and spinal cord (not shown). However, RT-PCR has shown the presence of this enzyme in testis (not shown).

A single RNA band of approximately 3.0 kb was detected with the specific probe for NL-3 (not shown). NL-3 expression was observed mainly in ovary, spinal cord and adrenal gland.

Chromosomal localisation of the human gene for NL-2 and NL-3

As a mean to get clues on the function of the new metallopeptidases in vertebrates, we have localized the new cDNAs on human chromosomes, in order to look for a possible link between the gene locus and mapped genetic diseases in humans. To do so, we have mapped the NL-2 and NL-3 genes by high-resolution fluorescence *in situ* hybridization (FISH). NL-2 was localized to chromosome band 1p36. Consistent with the cellular distribution of NL-2 in humans, genetic diseases of the CNS such as dyslexia, neural tube defect, neuroblastoma, neuronal type of Charcot-Marie-Tooth disease have all been mapped in this region and represent potential targets for a role of NL-2 in humans. NL-3 was localized to chromosome band

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2q37. Consistent with a role of NL-3 in bones, a form of Albright hereditary osteodystrophy was mapped to the same chromosomal locus (Phelan et al., 1995).

In view of the foregoing, NL-2 and NL-3 are metallopeptidases that are assumed to be immediately useful as markers for a disease or disorder associated with human chromosomal locus 1p36 and 2q37, respectively. Their localization on a chromosome band associated with known diseases suggests that they may be expressed or co-expressed with one or more genes, as a cause or a consequence of disease development. It is possible that these enzymes are up or down regulated, alone or along with other genes involved in a disease. Therefore, antibodies or other ligands specific to NL-2 or NL-3 may be used for a diagnostic purpose, as well as primers or probes in diagnostic assays using nucleic acid hybridization or amplification techniques. Otherwise, primers or probes directed against the nucleic acids of NL-2 and NL-3 would be useful to map the mutations of a gene located in close proximity and involved in the disease. Therefore, no matter which exact function NL-2 and NL-3 gene products have, their chromosomic localization provides one diagnostic utility. This localization as well as the tissular distribution provide information as to the disease and tissue to be investigated to elucidate the exact function of these enzymes.

NL-1 resembles NL-2, sharing with the latter about 80% homology in the amino sequence and sharing structural characteristics such as the furin recognition sequence located at the proximal end of the ectodomain. NL-2 might be the human homologue of mouse NL-1. If such was the case, these two proteins would have a substantial degree of divergence and, maybe, different profiles of activity varying from one species to another.

Chromosomal localization of NL-1 was determined in mouse genome by Single Strand Conformational Polymorphism (SSCP) in collaboration with The Jackson Laboratory Backcross DNA Panel Mapping Resource. NL-1 was localized to the distal region of mouse chromosome 4 which corresponds to human chromosome region 1p36 where is located NL-2 gene. This reinforces our hypothesis that NL-1 and NL-2 are species variants.

Production of antibodies against NLs

Antisera collected from injected animals were first tested by immunoblotting on GST-antigen fusion proteins produced in *E. coli*. Antiserum from one rabbit recognized the NL-1-related polypeptide and antisera from one mouse and one rabbit reacted with the NL-3-related polypeptide (results not shown). The anti NL-1 antiserum and the mouse anti NL-3 antiserum, which appeared more specific than the rabbit antiserum, were next tested by immunoblotting on extracts of proteins and culture media from cells expressing NL-1 or NL-3 (see below).

Expression of NL-1 in CHO cells

The cDNA encoding the full-length NL-1 protein was cloned in the mammalian

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expression vector pcDNA3-RSV and transfected in CHO cells. Stable cell lines were established by selection with the drug G418 and tested by immunoblotting for the presence of NL-1.

Small amounts of NL-1 were found in the extracts of transfected CHO cells (results not shown). This intracellular species was sensitive to endo H digestion, indicating that the sugar moiety was not mature and suggesting ER localization (results not shown). The culture medium of transfected CHO cells showed the presence of soluble NL-1 (Figure 10). This extracellular species was resistant to endo H suggesting true transport through the late secretory pathway. The cDNA sequence of NL-1 predicts a type II transmembrane protein. The mechanism by which NL-1 is transformed into a soluble protein is not known presently. However, examination of the amino acid sequence revealed the presence of a putative furin cleavage site from residue 58 to 65 (Figure 3). A similar site is present in NL-2 sequence.

The soluble form of NL-1 was assayed for activity using [3 H]-Tyr-(D)Ala $_2$ -Leuenkephalin and bradykinin as substrates. Figure 11 shows that NL-1 can degrade the enkephalin substrate ($K_m = 18\pm10~\mu\text{M}$) and that this activity can be inhibited by phosphoramidon (IC_{50} =0.9±0.3 nM) and thiorphan (K_m =47±12nM), a general inhibitor of enzymes of the NEP family. Bradykinin is also a substrate for NL-1 (not shown). Use of NL-1 amino-terminal domain to promote secretion

The observation that NL-1 ectodomain was secreted, possibly through cleavage of the transmembrane segment by furin, raised the possibility to promote secretion of exogenous proteins that could be spliced to NL-1 amino-terminal domain (from initiator methionine to the furin site). To test this hypothesis, the ectodomain of NL-3 (from the third cysteine to the end) was spliced to NL-1 amino-terminal domain using a PCR strategy and the recombinant DNA cloned in expression vector pRcCMV. The fusion protein was expressed by transfection of the vector in COS-1 and HEK 293 cells. The culture media of transfected cells was analyzed by immunoblotting using the mouse antiserum against NL-3. Figure 12 shows the presence of NL-3 in the spent culture media of both COS-1 and HEK 293 cells. This result shows that NL-1 amino-terminal domain can be used to promote secretion of exogenous proteins.

The soluble form of NL-3 was assayed for activity using $[^3H]$ -Tyr-(D)Ala₂-Leuenkephalin as substrate. No activity was found.

The previous experiment showed that it was possible to use the amino-terminal domain of NL-1 to promote secretion of an otherwise membrane attached protein ectodomain. To verify whether the same strategy could be used to promote secretion of small peptides, a PCR strategy was used to splice human β-endorphin to the amino-terminal domain of NL-1 and the recombinant DNA was cloned in vector pRcCMV. The fusion protein was expressed by transfection of the vector in COS-1 and HEK 293 cells. The culture media of transfected cells was collected 48h after transfection and

the peptides purified as described previously (Noël et al., 1989). The presence of β -endorphin in the extracts was detected by radioimmunoassay. The results showed that both COS-1 and HEK 293 cells produced approximately 100 pg of β -endorphin per ml of culture medium. Therefore, the N-terminus of LN-1 and NL-2 which ends with a furin-recognition site will be useful to produce the soluble form of a protein of interest.

SECTION 2)

MATERIALS AND METHODS

10 DNA manipulations

All DNA manipulations, phage library screening, and plasmid preparations were performed according to standard protocols (Ausubel 1988; Sambrook 1989). Site-directed mutagenesis was performed using a PCR-based strategy as described previously (Le Moual 1994).

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mRNA purification and RT-PCR protocol for identification of new members of the neprilysin family

mRNAs were prepared from mouse testis using Quick Prep Micro mRNA purification kit (Pharmacia Biotech). First strand cDNA was synthesized from 1 µg of mRNA using the First-Strand cDNA synthesis kit (Pharmacia Biotech).

Two oligonucleotide 3817 (5'sense primers. TGGATGGAT/CGA/CIGG/AIACIA/CA-3') oligonucleotide 3719 (5'and A/GTIGTITTT/CCCIGCIGGIA/GT/AIC/TTG/CCA-3') corresponding respectively to amino acid residues 459 to 465 and 552 to 560 of NEP sequence, and one antisense primer, oligonucleotide 3720 (5'-AIICCICCIA/TC/TA/GTCIGCIG/AC/TA/GTTT/CTC-3') corresponding to amino acid residues 646 to 654 (see Fig. 1 and 2), were synthesized. PCR was performed with 5 µl of cDNA template and 1 µl of Taq DNA polymerase in a final volume of 100 µl, containing 1 mM MgCl₂, 2 µM of each oligonucleotide 3817 and 3720, 200 µM of each dNTP and 5% DMSO. Cycling profiles included an initial denaturation step of 5 min at 94°C, 30 cycles of 1 min at 94°C, 1 min at 40°C and 1.5 min at 72°C, and a final extension step at 72°C for 10 min. One half of the amplified DNA was fractionated on a 2% agarose gel and fragments ranging in size between 500-700 bp were purified and resuspended in a final volume of 50 µl. A second round of PCR was done with primers 3719 and 3720, using as template either 10 µl of the first PCR reaction or 5 µl of the purified fragments, and the new PCR products were ligated in pCR2.1 vector (Invitrogen). Several identical clones corresponded to a potential new member of the NEP family. We called this member NL1 for NEP-like 1.

Cloning of full-length NL1 cDNA

The cloned NL1 PCR fragment was used as probe to screen a mouse testis λ Uni-ZAP™XR cDNA library (Stratagene). Twelve out of a hundred positive phages were plaque purified and subcloned into pBS SK vector (Stratagene). As the longest clone analyzed presented an incomplete ORF (pBS-NL1A), 5'RACE with primers located in vector (5'-TAGTGGATCCCCGGGCTGCAG-3', sense primer) and NL1 (5'-ACCAAACCTTTCCTGTAGCTCC-3', antisense primer, nt 1303 to 1324 of NL1; was subsequently performed on the DNA of the remaining semi-purified positive clones. Amplification was performed with 1µl of Vent polymerase in a final volume of 100 µl containing 50 ng of DNA, 4 mM of MgSO₄, 1µM of each oligonucleotide, 200 µM of each dNTP and 10% DMSO. Cycling parameters included an initial denaturation step of 1 min at 94°C, 25 cycles of 30 sec at 94°C, 30 sec at 60°C and 1 min at 72°C, and an incubation of 10 min at 72°C. A PCR fragment of the expected length was subcloned into pCR2.1 vector (clone pCR-NL1A), but sequencing revealed no initiator ATG codon. A nested 5'RACE was then performed on mouse testis cDNA using the Marathon Ready cDNA kit (Clontech) with sense oligonucleotides AP1 and AP2 (from the kit) and NL₁ antisense oligonucleotides 5'-CCTGAGGGCTCGTTTTACAACCGTCCT-3' (nt 503 to 529 of NL1) and 5'-CTCATCCCAGGAGAAGTGTAGCAGGCT-3' (nt 475 to 502 of NL1) as recommended by the supplier. The resulting fragment was cloned into pCR2.1 vector (pCR-NL1B). Since only ten bp were missing for the initiator ATG codon, we reconstructed the 5' end of the cDNA by PCR-amplifying clone pCR-NL1A with sense primer 5'-CCACCATGGTGGAGAGAGCAGGCTGGTGTCGGAAGAAG-3' (nt 332 to 364 of NL1; the 10 missing nucleotides are underlined) and antisense primer 5'-ACCAAACCTTTCCTGTAGCTCC-3' (nt 1303 to 1324 of NL1) using Vent polymerase as described above. The DNA fragment was then inserted into pCR2.1 (clone pCR-NL1C). The entire ORF was reconstituted following digestion of pBS-NL1A and pCR-NL1C with EcoRI and PfIMI. The 5' end of NL1 cDNA was excised from pCR-NL1C and ligated into pBS-NL1A at the corresponding sites, resulting in plasmid pBS-NL1B.

For expression studies, a *BamHI/ApaI* fragment generated out of pBS-NL1B, corresponding to the full length cDNA of NL1, was inserted into pCDNA3/RSV [18] vector.

Production of polyclonal antibodies

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A plasmid for the production in *Escherichia coli* of a GST fusion protein with NL1 was constructed using pGEX-4T-3 expression vector (Pharmacia Biotechnologies). A 255 bp fragment from NL1 was amplified by PCR with Vent polymerase using sense primer 5'-GCTACGGGATCCGTGGCCACTATGCTTAGGAA-3' (nt 1139 to 1158) and antisense primer 5'-CGATTGCTCGAGTGGGAACAGCTCGACTTCCA-3' (nt 1377 to 1396). Both pGEX-4T-3 and the PCR product were digested with *Bam*HI and *Xho*I

and ligated. The recombinant protein was produced and purified according to the supplier's instructions. Five weeks old female balb/c mice were immunized at monthly intervals for 3 months with 20 µg of the recombinant NL1 fragment in Freund's adjuvant and antisera were subsequently collected.

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Cell culture and transfection

Human Embryonic Kidney (HEK 293) cells were maintained in DMEM medium containing 10% fetal bovine serum (FBS), and supplemented with penicillin at 60 μ g/ml, streptomycin at 100 μ g/ml and fongizone at 0,25 μ g/ml. Transfections of cells with appropriate plasmids were performed by the calcium/phosphate-DNA coprecipitation method (Chang 1987). To establish permanent cell lines, G418 selection was initiated 48 h after the transfections at 400 μ g/ml for 12 days and gradually decreased at 100 μ g/ml.

LLC-PK₁ cells transfected with pRcCMV-sNEP were maintained as described previously (Lanctöt 1995).

Immunoblot analysis

For immunoblot analysis, cells were incubated for 16 h in synthetic DMEM medium containing 2mM sodium butyrate. Cellular proteins were solubilized as previously described (Dion 1995). Secreted proteins recovered in culture media were concentrated approximately 10 fold by ultrafiltration. Immunoblot analysis were performed using the NEN Renaissance kit with the polyclonal antibody specific to NL1 or the α 1-antitrypsin inhibitor antibody (Calbiochem; LaJolla, CA) followed by the appropriate horseradish peroxidase-conjugated IgG (Vector Laboratories).

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For the glycosylation studies, proteins were incubated with endoglycosidase H (endoH) or peptide:N-glycosidase (PNGaseF) as suggested by the distributor (NEB).

Enzymatic activity assays

NL1 activity was monitored and compared to sNEP activity using (Tyrosyl-[3,5- 3 H])(D-Ala₂)-Leu₅-enkephalin (50 Ci/mmol) (Research Products International Inc.), as already described (Dion 1995; Devault 1988). K_m values were determined by the isotope-dilution method. The inhibitory effects of phosphoramidon and thiorphan were also assessed as previously described (Dion 1995).

35 HPLC analysis of the hydrolysis of Leu-enkephalin

Five μg of Leu_s-enkephalin were incubated at 37°C for one hour in 50mM MES, pH 6.5, with concentrated culture medium of HEK 293 cells expressing NL1 (~300 μg of total proteins) or LLC-PK₁ cells expressing sNEP (~30 μg of total proteins), in absence or presence of 0.1 mM phosphoramidon. Hydrolysis products were separated by

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reversed-phase HPLC as described previously [23]. Tyr-Gly-Gly and Phe-Leu were both identified by elution profiles of synthetic marker peptides.

Northern blot analysis

A mouse multiple tissue poly(A)⁺ mRNA blot (Clontech) was hybridized with a [³²P]dCTP random primer labelled probe in ExpressHyb solution (Clontech). The blot was washed according to the manufacturer's recommendations and exposed to Fuji RX film for 7 days at -80°C with intensifying screens.

10 RT-PCR screening of mouse tissues

First strand cDNA synthesis was performed with 1 µg of total RNA from mouse tissues and oligo(dT) as primer, using Gene Amp RNA PCR Core Kit (Perkin Elmer). For the PCR reactions, primers 5'-TGGCGAGAGTGTCAGCTATGTC-3' and 5'-CTTCCAAAATGTAGTCAGGGTAGCCAATC-3' were used with Taq polymerase. One tenth of the PCR products were visualized on a 4% agarose gel.

In situ hybridization

To construct a plasmid for the synthesis of cRNA probes for ISH, pCR-NL1A was used as template to amplify a 452 bp fragment by PCR with sense primer 5'-GGAGCCATAGTGACTCTGGGTGTC-3' (nt 416 to 439) and antisense primer 5'-GACGCTCAGCAGGGGCTCAGAGTC-3' (nt 842 to 865). The amplification product was inserted into pCRII vector (Invitrogen). Synthesis of riboprobes and protocols for ISH were as described previously (Ruchon 1998).

RESULTS

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Cloning and sequence analysis of mouse NL1 cDNA

In order to isolate cDNAs for new members of the NEP family, we developed an RT-PCR strategy based on fact that NEP, ECE-1 and PHEX share regions of significant sequence identity. Following RT-PCR on testis mRNAs with nested primers, a DNA fragment of approximately 300 bp was amplified. This DNA fragment was cloned and the plasmids from 24 independent colonies were sequenced: 3 clones had no insert, 4 clones had DNA fragments not related to the NEP family, 7 clones had sequences corresponding to mouse NEP and 3 clones had sequences corresponding to mouse PHEX, showing that our approach efficiently allowed the identification of members of the family. Moreover, 7 identical clones had a new cDNA presenting sequence similarities to members of the NEP family. The full-length cDNA was subsequently obtained by screening a mouse testis λ cDNA library followed by 5'RACE, as described under *Materials and Methods*. Its nucleotide and deduced amino acid sequences confirm that we cloned a novel NEP-like protein, referred to

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thereafter as NL1.

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NL1 cDNA spans 2925 nt, including a 5'-untranslated region of 331 nt, an open reading frame of 2295 nt from nt 332 to nt 2626, and a 3'-untranslated region of 299 nt. The sequence surrounding the proposed initiator ATG conforms to the Kozak consensus (Kozak 1986). The deduced amino acid sequence of NL1 reveals a putative type II transmembrane protein of 765 amino acid residues encompassing a short N-terminal cytoplasmic tail, a unique putative transmembrane domain, and a large C-terminal extracellular domain. The ectodomain contains nine potential N-glycosylation sites (Asn-X-Ser/Thr) and ten cysteine residues corresponding to those conserved among all the members of the family, which are presumably involved in proper folding and in maintenance of the protein in an active conformation. All amino acid residues known to be part of the active site of NEP are present in NL1. The predicted protein presents greater similarities to NEP than to any other member of the family.

Although NL1 shares numerous features with proteins of the neprilysin family, a notable aspect distinguishes it from the others: the first conserved cysteine residue of the ectodomain is more distant (34 amino acid residues) from the predicted transmembrane domain in NL1 than it is in NEP (9 residues) or any other members of the family. Moreover, we noticed a putative furin cleavage site (-Arg₅₈-Thr-Val-Val-Lys-Arg₆₃-) between the end of the transmembrane domain and the first cysteine. This observation suggests that NL1 could exist as a secreted rather than a membrane-bound protein.

NL1 expression in HEK 293 cells

HEK 293 cells were transfected with pCDNA3/RSV expression vector containing NL1 cDNA, and a permanent cell line was established as described under Materials and Methods (HEK/NL1 cells). Immunoblotting with a polyclonal antibody showed that after 16h of culture, most NL1 was present in the culture medium with small amounts of the enzyme in the cell extract. Secreted and cell-associated NL1 had apparent molecular masses of approximately 125 and 110 kDa, respectively. To characterize the glycosylation state of NL1, we next submitted the recombinant protein to deglycosylation by peptide: N-glycosidase F (PNGase F) and endoglycosidase H (endo H). PNGase F removes high mannose as well as most complex N-linked oligosaccharides added in the Golgi complex. In contrast, endo H removes N-linked oligosaccharide side chains of the high mannose type found on proteins in the RER but which have not yet transited through the Golgi complex; thus, resistance to endo H can be used as an indication that the protein has traveled through the Golgi complex. PNGase F treatment showed that the cell-associated and secreted NL1 were N-glycosylated as their electrophoretic mobility increased following digestion. However, the secreted NL1 migrated as a doublet after PNGase F treatment, with one

band co-migrating with cell-associated form and the second having a slower rate of migration. Since untreated and endo H-digested secreted NL1 are seen as single bands by SDS-PAGE, our observation suggests that a proportion of secreted NL1 undergoes further post-RER postranslational modification that renders some of the N-

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linked oligosaccharides resistant to PNGase F digestion.

In contrast to secreted NL1, NL1 from cell extract was sensitive to endo H treatment. This result shows differences in the glycosylation state of the two species and suggests that the cell-associated form observed in transfected cells is an intracellular species that has not traveled through the Golgi complex.

Processing of NL1 by a subtilisin-like convertase

To determine whether a member of the mammalian subtilisine-like convertase family is responsible for NL1 presence in the culture medium of transfected cells, we co-transfected transiently HEK 293 cells with a constant amount of plasmid pCDNA3/RSV/NL1 and increasing amounts of plasmid pCDNA3/CMV/PDX (Benjannet 1997). This latter vector promotes the expression of the α 1-antitrypsin Portland variant, α 1-PDX, a known inhibitor of subtilisin-like convertases (Anderson 1993). Immunoblot analysis of the culture media of cells expressing both NL1 and α 1-PDX indicated that NL1 secretion was strongly inhibited by the presence of α 1-PDX: a relation was observed between the amounts of α 1-PDX and the level of inhibition of NL1 secretion.

To confirm that proteolysis by the subtilisin-like convertase occurred at the putative furin cleavage site identified in NL1 ectodomain (-Arg₅₈-Thr-Val-Val-Lys-Arg₆₃-), the amino acid residues Asn₆₂-Gly₆₃ were substituted for Lys₆₂-Arg₆₃ by site-directed mutagenesis in vector pCDNA3/RSV/NL1 and the mutated vector used to establish HEK 293 cells expressing the mutant protein (HEK/NL1mut cells). Immunoblot analysis of the culture media of HEK/NL1mut cells showed that the mutation totally abolished secretion of NL1. Furthermore, an additional form of NL1 with a molecular mass of 127 kDa was detected in the extract of these cells. This new species was resistant to endo H digestion and was found associated with membranes when HEK/NL1mut cells were fractionated according to Chidiac *et al.* 1996 (result not shown).

NL1 enzymatic activity

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Culture media from HEK 293 and HEK/NL1 cells were tested for enzymatic activity using as substrate (Tyrosyl-[3,5-³H])(D-Ala₂)-Leu₅-enkephalin, a known NEP substrate. Activity was detected in the culture medium of HEK/NL1 cells but not in that of HEK 293 cells. This activity increased linearly with the amounts of NL1 and with the incubation period, indicating that degradation of the substrate was due to NL1 enzymatic activity.

We next characterized NL1 enzymatic parameters using the same substrate

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and compared them to those of an engineered soluble form of NEP (sNEP) (Lemay 1989). NL1 affinity for D-Ala₂-Leu₅-enkephalin was slightly higher than that of sNEP as shown by their K_m values of 18 μ M and 73 μ M, respectively. Inhibition assays showed that phosphoramidon had similar effects on NL1 and sNEP activity, with IC₅₀ values of 0.9 nM and 0.5 nM respectively, and that thiorphan, a specific inhibitor of NEP, inhibited NL1 with an IC₅₀ of 47 nM, as compared with an IC₅₀ of 8 nM for NEP.

Very low levels of phosphoramidon-sensitive activity was detected in extracts of HEK/NL1 cells (data not shown) consistent with the small amounts of NL1 observed by immunoblotting.

To determine whether NL1 had cleavage site specificity similar to NEP, we incubated Leu $_5$ -enkephalin in the presence of NL1 recovered from the medium of HEK/NL1 cells or in the presence of sNEP, and analyzed the degradation products by RP-HPLC. Peaks co-migrating with standard Tyr-Gly-Gly and Phe-Leu peptides were observed in both RP-HPLC profiles, indicating that both enzymes cleaved the substrate at the Gly $_3$ -Phe $_4$ peptide bond. This enkephalin-degrading activity was completely inhibited by phosphoramidon (1 μ M).

Tissue and cellular distribution of NL1 mRNA

Tissue distribution of NL1 mRNA was determined by Northern blot analysis with a specific probe corresponding to the 5'end of the coding region of NL1 cDNA. A single transcript of 3.4 kb was detected exclusively in testis among all the mouse tissues tested. Mouse tissues were also screened by RT-PCR. Using this more sensitive technique, expression of NL1 was observed in several other tissues including heart, brain, spleen, lungs, liver and kidney. Consistent with the Northern blot results, RT-PCR analysis, although not strictly quantitative, detected more NL1 mRNA in testis than in other tissues.

To gain more insight into NL1 mRNA distribution, we examined by ISH cryostat sagital sections from a 4-day newborn mouse, as well as sections from a 16-day old animal (p16) and adult tissues (heart, brain, spleen, lungs, liver, kidney and testis). The presence of NL1 mRNA was detected only in adult testis. Only the germinal cells in the luminal face of the seminiferous tubules were labeled. These cells were identified as round and elongated spermatids in all spermiogenesis maturational stages. Neither spermatozoa nor spermatocytes, spermatogonies or Sertoli cells were labeled. Interstitial cells were also negative. Controls were performed with sense riboprobes, which produced only nonspecific background (data not shown). The 4-day old mouse sagital sections and all other tissues tested were negative.

DISCUSSION

The great interest in members of the Neprilysin family as putative therapeutic

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targets, and the recent discovery of new members of this important family of peptidases led us to investigate whether additional members of the family remained to be identified. Using a PCR-based strategy, we cloned, from mouse testis, a partial cDNA encoding a new NEP-like enzyme that we called NL1. Analysis of the amino acid sequence encoded by the full-length NL1 cDNA revealed that this member of the family resembles NEP the most: 55% identity and 74% similarity. Recently, the primary structure of a new zinc metallopeptidase from total mouse embryo was reported (Ikeda 1999). This enzyme, called SEP, is found either as a soluble or a cell-associated form due to alternative splicing. NL1 shows only 3 amino acid differences with the soluble form of SEP indicating that secreted SEP and NL1 are the same enzyme. Our cloning strategy did not allow characterization of the cell-associated form of NL1 which is a minor species in mouse testis (Ikeda 1999).

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The amino acid sequence of NL1 predicts a topology of a type II integral membrane glycoprotein that is similar to the other members of the family. Treatment of the recombinant protein with PNGase F showed that indeed NL1 possesses N-linked carbohydrate side chains. However, it is not possible to determine precisely whether all nine putative N-glycosylation sites are used, but the 30 kDa decrease in molecular mass upon PNGase F treatment suggests that most are glycosylated. It has already been shown that all asparagine residues in a Asn-X-Ser/Thr consensus are glycosylated in rabbit NEP expressed in COS-1 cells and that sugar moieties increase the stability and enzymatic activity of the protein and facilitate its intracellular transport (Lafrance 1994). Three of NEP glycosylated Asn residues (Asn 145, Asn 285 and Asn 628) are conserved in NL1 (Asn 163, Asn 303 and Asn 643). Amongst these residues, Asn 145 and Asn 628 have been reported to influence NEP enzymatic activity (Lafrance 1994). In the same work, it has also been shown that the effect of sugar addition on folding and intracellular transport of NEP is a cumulative effect of all glycosylation sites rather than a contribution of any particular one. Glycosylation of NL1 may share similarities with that of NEP since we found their primary structures and enzymatic activities to be very similar.

Surprisingly, expression of the cDNA by transfection of HEK 293 cells showed that most of the enzyme was secreted in the culture medium. The small amount of NL1 associated with the cells was endo H-sensitive, suggesting that the cell-associated enzyme is a species that has not yet left the RER. The presence of a furin cleavage site in NL1 sequence between the predicted transmembrane domain and the first conserved cysteine residue of the ectodomain led us to believe that a member of the mammalian subtilisin-like family of convertases was responsible for the presence of NL1 in the culture medium. These enzymes are involved in processing a variety of precursor proteins such as growth factors and hormones, receptors, plasma proteins, matrix metalloproteinases, metalloproteases-desintegrins and viral envelope

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glycoproteins [for a review see: (Nakayama 1997). Site-directed mutagenesis of the furin cleavage site (-Arg₅₈-Thr-Val-Val-Lys-Arg₆₃-) and expression of α 1-PDX, a potent inhibitor of mammalian subtilisin-like convertases (Anderson 1993), confirmed that a member of this family of endoproteases was involved in NL1 secretion presumably by cleaving in carboxy-terminus of Arg₆₃. There are only a few examples of proteins which are processed from a membrane-bound precursor to a secreted form following cleavage by subtilisin-like convertases; these include meprin and collagen XVII (Milhiet 1995; Schacke 1998). Three members of the subtilisin-like family of convertases, namely furin, PC4 and PC7, are known to be expressed in germ cells (Nakayama 1992; Torri 1993; Seidah 1992, 1996). Whether one of these convertases generates secreted NL1 from its membrane form is under current investigation. In any case, NL1 is the only known member of the neprilysin family that is secreted. This unique feature suggests that NL1 plays its physiological role in a context different from that of the membrane-bound peptidases, thereby diversifying the role of the peptidases of the neprilysin family. It is of interest that circulating forms of NEP in blood and urine have been described, but they have generally been related to pathological or stressful conditions (Almenoff 1984; Deschodt-Lanckmann 1989; Johnson 1985; Soleilhac 1996; Aviv 1995).

We have observed in cells expressing NL1 mutated at the furin cleavage site the appearance of a species resistant to digestion by endo H. This mutated protein was associated with cellular membranes. Taken together, these results indicate that NL1 is first synthesized and inserted in the RER membrane as a type II transmembrane protein. During intracellular transport, NL1 is converted to a soluble form by the action of a member of the mammalian subtilisin-like convertases. The identity of the cellular compartment where this process occurs is not known. However, mammalian subtilisin-like convertases are usually active in post-Golgi compartments of the secretory pathway suggesting that processing of NL1 from the membrane bound form to the soluble form is a post-Golgi event.

Despite almost total abrogation of NL1 secretion, we observed only a slight accumulation of endo H-resistant NL1 in cells either co-expressing α 1-PDX and NL1 (result not shown) or expressing mutated NL1. This observation suggests that unprocessed NL1 is rapidly degraded. A similar behavior was reported for the Notch1 receptor expressed in the furin-deficient cell line LoVo (Logeat 1998). The mechanism(s) by which these unprocessed proteins are degraded is still unknown. It is interesting to point out that the spliceoform of SEP that has lost a 23 amino acid peptide, including the furin cleavage site, generates a cell-associated endo H-sensitive molecule (Ikeda 1999).

The most important observation regarding the NL1 primary structure is the conservation of residues which in NEP are essential for catalysis and binding of

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substrates or inhibitors. This finding suggests that NL1 could effectively act as an endopeptidase with a catalytic mechanism similar to that of NEP. This hypothesis was supported by the demonstration that D-Ala₂-Leu₅-enkephalin, a peptide substrate often used to monitor NEP activity, was also an excellent NL1 substrate. The affinity of NL1 for D-Ala₂-Leu₅-enkephalin was even higher than that of NEP, as reflected by a K value 4- to 5-fold lower. Furthermore, two well known NEP inhibitors, phosphoramidon and thiorphan, also abolished NL1 activity. Phosphoramidon, which inhibits NEP as well as ECE-1 activity, albeit to a lesser extent (Turner 1996), had very similar effects on NL1 and NEP, with an IC₅₀ value for NL1 varying not more than two-fold from the value determined for NEP. Thiorphan, considered to be a more specific inhibitor of NEP, also inhibited NL1 activity, with an IC_{50} six-fold greater than that for NEP. These results suggest that the active sites of NL1 and NEP are similar. This hypothesis is supported by the observation that secreted SEP degraded a set of peptides known to be NEP substrates, including substance P, bradykinin and atrial natiuretic peptide (Ikeda 1999). Taken together, these results illustrate the importance of identifying and characterizing other member of the family for the design of highly specific inhibitors.

In agreement with the enzymatic parameters demonstrating that NL1 and NEP have similar catalytic sites, we have observed that both enzymes cleaved Leu₅-enkephalin at the same peptide bond. This result suggests that NL1 hydrolyzes peptide bonds on the amino side of hydrophobic amino acid residues as does NEP (Turner 1985). However, several other peptides will have to be tested to confirm this specificity and to determine whether NL1 has dipeptidyl carboxypeptidase activity as was shown for NEP (Malfroy 1982; Bateman 1989; Beaumont 1991) and more recently for ECE-1 (Johnson 1999).

RT-PCR experiments with specific primers for the soluble and cell-associated forms of SEP showed a wide tissue distribution of the enzyme with the soluble form of SEP being predominant in testis and the cell-associated form in other tissues (Ikeda 1999). Our RT-PCR results confirmed the wide tissue distribution of NL1. However, Northern blotting and *in situ* hybridization experiments indicated that expression of NL1 is predominant in germ cells of mature testis. Interestingly, proenkephalin mRNA has been shown to be expressed in germ cells and somatic cells of the testis (Torii 1993, Seidah 1992; Kew 1989; Mehta 1994; Kilpatrick 1986, 1987). Specific functions for testicular enkephalin peptides have not yet been defined, but it is believed that they could act as intratesticular paracrine/autocrine factors. In addition to their putative role as mediators of testicular cell communication, it has also been demonstrated that proenkephalin products synthesized by spermatogenic cells during spermatogenesis are stored in the acrosome of human, hamster, rat and sheep spermatozoa and are release from sperm following acrosomal reaction (Kew 1990). It has thus been proposed that proenkephalin products may act as sperm acrosomal factors during the

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fertilization process as well as intratesticular regulators secreted by spermatogenic cells. Since Leu₅-enkephalin was found to be a good substrate for NL1, opioid peptides originating from proenkephalin could serve as physiological substrate for this new enzyme. In this way, NL1 would serve to regulate the activity of these bioactive peptides.

Testis is the only tissue where the soluble form of SEP is predominant (Ikeda, 1999), suggesting a testis-specific alternative splicing. Expression of testis-specific molecular species of peptidases or prohormones, arising through diverse mechanisms, has been documented in the past (Howard 1990; Jeannotte 1987). However, the physiological significance of these testis-specific species is not always clear. In the case of NL1 or SEP, it might allow local constitutive secretion by germinal cells of an otherwise cell-associated enzyme, to regulate spermatogenesis much like several other proteolytic enzymes of the seminiferous tubules (Monsees 1998). Alternatively, it might allow accumulation in acrosome with proenkephalin peptides and release upon acrosomal reaction. More exhaustive studies concerning NL1 localization and physiological substrate identification will be needed to understand its role in the testis and possibly in the fertilization process.

Cloning of other members of the family

To find other members of the NEP-like family, we will use the same RT-PCR strategy to amplify mRNA isolated from tissues known to be regulated by peptidergic systems (brain, thymus, kidney, heart, lung, ovary, pancreas, bone, bone marrow and lymphoid cells). In fact, many of these tissues are known to express at least one member of the family and/or to control a peptidergic pathway on which peptidase inhibitors have major effects. Amplified fragments will be cloned and the resulting clones will be sequenced and compared to the sequence of known peptidases, as described above. Pairs of degenerate primers in other highly conserved regions will also be designed to increase the possibility of cloning other relevant peptidases.

DISCUSSION

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As discussed above, peptidases of the NEP family known to date have often been found to play important physiological roles. This is certainly true for NEP itself, ECEs and PEX, (see review above). For this reason, some of these enzymes (as it was the case for NEP and ECE in the past) might be interesting targets for the design of inhibitors that in turn could be used as therapeutic agents in various pathological conditions. However, it is of some concern that inhibitors designed for one enzyme may also inhibit to some extent other members of the family. This lack of specificity for an inhibitor used as a therapeutic agent in the long term treatments such as those used as antihypertensive agents for instance, may cause unforeseen problems due to unwanted side effects. The objectives of the present work was to develop a strategy

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to clone new members of the NEP family of peptidases. The results presented in this report clearly show that our strategy can be successful. We have determined the complete or partial nucleotide sequence of three cDNAs encoding putative enzymes of the NEP family.

These cDNA sequences are valuable tools and may be used to: Produce antibodies

As shown in the present work, knowledge of NL cDNA sequences can be used to raise specific antibodies. For example but not exclusively, regions of less homology between the peptidases (amino acid residues 50 to 450) can be used to synthesize peptides whose sequences are deduced from the translation of the cDNAs, and/or bacterially-expressed fragments of the cDNAs fused for example but not exclusively to GST may be purified and injected into rabbits or mice for polyclonal or monoclonal antibody production. These antibodies can be used to:

- identify by immunohistochemistry the peptidergic pathways in which the peptidases are functioning;
- study the physiopathology of NL-enzymes by immunoblotting or immunohistochemistry on samples of biological fluids or biopsies;
- set up high through put screening assays to identify NL-enzymes inhibitors. This can be done for example but not exclusively by using the antibodies to attach the NL-enzymes to a solid support;
- purify NL-enzymes with said antibodies by immunoprecipitation or affinity chromatography by identifying antibodies capable of selectively binding to the NL-enzymes in one set of conditions and releasing it in another set of conditions typically involving a large pH or salt concentration change without denaturing the NL-enzyme;
- identify antibodies that block NL-enzymes activities and use them as therapeutic agents. Blocking antibodies can be identified by adding antisera or ascite fluid to an *in vitro* enzymatic assay and looking for inhibition of NL-enzymes activities. Blocking antibodies could then be injected to normal or disease model animals to test for *in vivo* effects.

Derive specific RNA or DNA probes

As shown in the present work, knowledge of the nucleotide sequence of the members of the NEP-family allows nucleotide sequence comparisons and facilitate the design of specific RNA or DNA probes by methods such as but not exclusively molecular cloning, *in vitro* transcription, PCR or DNA synthesis. The probes thus obtained can be used to:

derive specific probes or oligonucleotides for RNA and DNA analysis, such as Northern blot and *in situ* hybridization, chromosome mapping

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or PCR testing. These probes could be used for genetic testing of normal or pathological samples of biological fluids or biopsies;

make vectors for gene knock-out or knock-in in mice. The long range PCR technique and/or screening of a mouse genomic library with probes derived from the 5'-end of the cDNAs can be used to isolate large exon/intron regions. We will then substitute one or more of the cloned genomic DNA exons for the neomycin resistance expression cassette for producing homologous recombination and knock-out mice. Alternatively, cDNAs coding for NLs will be used to overexpressed each of these enzymes in transgenic mice. The cDNAs will be cloned downstream from a promoter sequence, and injected in fertilised mouse eggs. Depending on specific questions to be answered, the chosen promoter sequence will allow expression of the peptidases either in every tissues or in a cell- or tissue-specific manner. Injected eggs will be transferred into foster mothers and the resulting mice analysed for peptidase expression;

replace defective NL genes in a gene therapy strategy. The NL full length cDNAs could be cloned under the control of a constitutive or inducible promoter having a narrow or wide range of tissue expression and introduced with appropriate vectors in subjects having defective genes;

synthesise oligonucleotides that could be used to interfere with the expression of the NLs. For example but not exclusively, oligonucleotides with antisens or ribozyme activity could be developed. These oligonucleotides could be introduced in subjects as described above;

isolate other members of the family. Screening cDNA and/or genomic libraries with these cDNA probes at low stringency may allow to clone new members of the NEP-like family. Alternatively, alignment of the sequences may allow one to design specific degenerate oligonucleotide primers for RT-PCR screening with mRNA from tissues such as but not exclusively, the hearth and the brain.

Production of recombinant NL-enzymes

As shown in the present work, recombinant active NL-enzymes can be obtained by expression of NL-cDNAs in mammalian cells. From past experience with neprilysin, another member of the family (Devault *et al.*, 1988; Fossiez *et al.*, 1992; Ellefsen *et al.*, submitted), expression can also be performed in other expression systems after cloning of NL-cDNAs in appropriate expression vectors. These expression systems may include but not exclusively the baculovirus/insect cells or larvae system and the *Pichia*

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pastoris-based yeast system. Production of recombinant NL-enzymes includes the production of naturally occurring membrane bound or soluble forms of the proteins or genetically engineered soluble forms of the enzymes. The latter can be obtained by substituting the cytosolic and trans-membrane domain by a cleavable signal peptide such as that of proopiomelanocortin, but not exclusively, as done previously (Lemay et al., 1989) or by transforming by genetic manipulations the non-cleavable signal peptide membrane anchor domain into a cleavable signal peptide, as done previously (Lemire et al., 1997) or by fusion of the ectodomain of NL-enzymes to the amino-terminal domain (from the initiator methionine to amino acid residue 300) of naturally occurring soluble NLs such as, but not exclusively, NL-1 as done in this work.

These recombinant NLs could be used to:

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- find a substrate. A substrate can be identified using one of the following.
- Screening of existing bioactive peptides. Peptides are incubated in the
 presence of NL-enzymes and subsequently analysed by HPLC for
 degradation. Degradation is observed by disappearance of the peak of
 substrate and the appearance of peaks of products;
- Screening phage libraries specifically designed for the purpose (phage display library). Each phage expresses at its surface, as part of its coat protein, a random peptide sequence preceded by a peptide sequence recognisable by an antibody or any other sequence-recognizing agent. This latter sequence serves to attach the phage to a solid support. Upon addition of the NL-enzyme the random sequences that are NL substrate are cleaved, releasing the phage. After several rounds of cleavage, the phage sequence is determined to identify the peptide segment recognized by the enzyme.
- Extract of the tissue where the enzyme is expressed is collected and prepared for chromatographic analysis (HPLC, capillary electrophoresis or any other high resolution separation system) by denaturing the extracted proteins with a solvent (acetonitrile or methanol). The extract is subjected to chromatographic separation. The same extract is incubated with the enzyme for a period sufficient to observe a difference between the 2 chromatograms. The regions with the identified changes are collected and subjected to mass spectrometric analysis to determine the peptide compositions.
- Small peptide libraries are prepared with a fluorophore at one extremity and a quencher group at the other (Meldal et al Methods in molecular biology 1998,87). The substrate can be identified using a strategy described in Apletalina et al (JBC (1998)273, 41, 26589-95). For each

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hexapeptide library, the identity of one residue at one position remains constant while the rest is randomized (for a total of 6*20=120 individual libraries). Each library is made-up of 3.2 million different members and is identified both by the position of the constant residue along the hexapeptide, and its identity. The NL-enzyme is added to each library and the fluorescence is recorded. The data is organized to identify the libraries producing the most fluorescence for each position along the hexapeptide. This arrangement suggests the identity of important residues at each position along the hexapeptide. Hexapeptide representing the best suggestions are prepared and tested in a similar fashion. From this set, the hexapeptide with the best fluorescence is selected.

set up enzymatic assays. An enzymatic assay consists in the addition of the above-identified substrate to the enzyme in constant conditions of pH, salts, temperature and time. The resulting solution is assayed for the hydrolysed peptide or for the intact peptide. This assay can be realized with specific antibodies, HPLC or, when self-quenched fluorescence tagged peptides are used (Meldal et al), by the appearance of fluorescence. The enzyme may be in solution or attached to a solid substrate;

identify inhibitors. Inhibitors can be identified from synthetic libraries, biota extracts and from rationally designed inhibitors using X-ray crystallography and substituent activity relationships. Each molecule or extract fraction is tested for inhibitory activity using the enzymatic test described above. The molecule responsible for the largest inhibition is further tested to determine its pharmacological and toxicological properties following known procedures. The inhibitor with the best distribution, pharmacological action combined with low toxicity will be selected for drug manufacturing. Pharmaceutically acceptable formulation of the inhibitor or its acceptable salt will be prepared by mixing with known excipients to produce tablets, capsules or injectable solutions. Between 1 and 500mg of the drug is administered to the patients;

inject the native or soluble purified NL-enzymes into subjects. In the case of disease or pathologies caused by a lack or decrease in NL activity, the purified NL could be injected intravenously or otherwise in patients. Alternatively, immobilized NL-enzymes could be introduced at the site of orthopedic surgery or implantation of devices in bones or dental tissues.

Secretion of foreign proteins and peptides

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As shown in the present work, the amino-terminal domain of NL-1 (from the initiator methionine to the furin site) can be used to promote the secretion of a foreign protein (in this case the ectodomain of NL-3 and β -endorphin).

The amino-terminal domain of NL-1 but also of other naturally occurring soluble NL-enzymes could be used to:

- promote production and secretion of foreign proteins. This can be achieved by genetically fusing sequences coding for said foreign proteins downstream from and in phase with the amino-terminal of NL-1. These chimeric constructs could be introduced with the help of appropriate vectors in any of the expression systems mentioned above for protein production and secretion;
- promote production and secretion of bioactive peptides. Sequences encoding small bioactive peptides such as but not exclusively β-endorphin, the enkephalins, substance P, atrial natriuretic peptide (ANF) and osteostatine, could be fused immediately downstream and in phase the furin site of NL-1. These DNA constructs could be used as described above to produce bioactive peptides.
 - serve as model to design artificial (non-naturally occurring) proteins or protein segments (protein vectors) to promote secretion of proteins or peptides. These protein vectors can be constructed to resemble a secreted protein. In this case they would be assembled of an endoplasmic reticulum signal peptide, a spacer of varying length and a furin cleavage site to which the protein or peptide destined for secretion can be fused. The total length of the spacer, furin cleavage site and protein or peptide destined for secretion must be at least 70 amino acid residues. Alternatively, such protein vectors could be assembled to resemble a type II membrane protein. In this case they would comprise from the amino to the carboxy-terminus a cytosolic domain of varying length, a transmembrane domain that also acts as a signal peptide, an extracellular segment of varying length and a furin cleavage site to which the protein or peptide destined for secretion can be fused. The total length of the extracellular segment, furin cleavage site and protein or peptide destined for secretion must be at least 70 amino acid residues.

Therapeutic applications of NL-enzymes

The inappropriate processing of endogenous peptides causes several diseases. The inappropriate processing may result from pathologic

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concentration of the enzyme itself, its substrate or other elements of the biochemical machinery downstream from the controlling enzyme. In this context it is possible to help the patient by managing the activity of the controlling enzyme.

NL-enzymes have been localized to the brain and may be involved in the improper processing of β-amyloid precursor. Inhibitions of this process by drugs prepared as above, will help patients with Alzheimer disease as well as other patient suffering from diseases caused by plaque formation;

 NL-enzymes may be involved in the improper processing of other peptides involved in neurological diseases, pain or psychiatric disorders.
 Appropriately designed inhibitors will help in the management of such diseases;

NL-1 is found in testis and is associated with spermatozoid maturation. Peptides improperly processed by the enzyme may lead to infertility. The addition of NL-1 ex-vivo to seminal liquid or immature spermatozoids taken directly from testis during an in-vitro fertilization procedure will increase fertility. Conversely, the use of a small-molecule inhibitor or removal of NL-1 with an antibody could increase fertility during an in-vitro fertilization procedure. The administration of a NL-1 inhibitor may increase or decrease the fertility potential. This inhibitor is formulated and administered as described above.

NL-3 is found in ovaries and may be involved in the processing of a peptide involved in the maturation of eggs. The addition of NL-3 ex-vivo to immature eggs taken directly from ovaries during an in-vitro fertilization procedure will increase fertility. Conversely, the use of a small-molecule inhibitor or removal of NL-3 with an antibody could increase fertility during an in-vitro fertilization procedure. This inhibitor is formulated and administered as described above;

NL-3 is found in bones. The improper processing of peptides by the enzyme may result in bone disease or abnormal phosphate metabolism. Administration of an inhibitor, as described above, will allow the disease management.

TABLE I

Extend of amino acid sequence identity between members of the NEP-like family

	hNEP	hPEX	hECE-1A	hECE-2	hKELL	sNL-1	hNL-2	hNL-3
hNEP	100*							

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hPEX	35	100						
hECE-1A	39	38	100					
hECE-2	36	37	62	100				
hKELL	23	24	30	31	100			
sNL-1	55	39	39	39	26	100		
hNL-2	54	39	39	39	26	77	100	
hNL-3	35	32	37	37	28	36	34	100

^{*:} percentage of sequence identity

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What is claimed is:

- 1. A neutral endopeptidase-like metallopeptidase which is found upon probing tissue nucleic acids with degenerate oligonucleotides derived from a conserved sequence located on either side of a sequence His-Glu-Xaa-Xaa-His, wherein Xaa is any amino acid, which has a sequence selected from the amino acid sequences shown in Figures 3, 4, and 5, a fragment thereof, and a variant thereof sharing at least about 80% homology with said sequence.
- 2. A metallopeptidase as defined in claim 1, which has the amino acid sequence shown in Figure 3.
 - 3. A metallopeptidase as defined in claim 1, which has the amino acid sequence shown in Figure 4.
- 4. A metallopeptidase as defined in claim 1, which has the amino acid sequence shown in Figure 5.
 - 5. A nucleic acid encoding the metallopeptidase of any one of claims 1 to 4.
- 20 6. A recombinant vector comprising the nucleic acid defined in claim 5.
 - 7. A recombinant host cell expressing the nucleic acid of claim 5.
- 8. A method for producing a metallopeptidase as defined in any one of claims 1 to 4, which comprises the steps of culturing a recombinant host cell as defined in claim 7 in a growth supportive medium and recovering said metallopeptidase from said host cell or the culture medium.
- 9. A method for screening new molecules related to neutral endopeptidase (NEP),30 which comprises the steps of:
 - aligning nucleotidic sequences of NEP and of known molecules related to NEP;
 - assessing consensus sequences on either side of a sequence comprising His-Glu-Xaa-Xaa-His sequence, wherein Xaa is any amino acid;
 - synthetising degenerate sequences of said consensus sequences;
 - contacting said degenerate sequences with the nucleic acids of a panel of samples susceptible to express said new molecules, in conditions such that a hybridization complex can form between the nucleic acids of samples and the degenerate sequences;

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- detecting the formation of said hybridization complex as an indication of a sample which comprises a molecule related to NEP; and
- sequencing the nucleic acid comprising said hybridization complex; whereby a new sequence sharing homology with NEP is a new molecule related to NEP.
- 10. A method as defined in claim 9, wherein said degenerate sequences are selected from Figure 2.
- 10 11. An oligonucleotide selected from those in Figure 2.
 - 12. A composition of matter comprising one or more of the oligonucleotides of claim 11.
- 15 13. A recombinant vector comprising a nucleic acid encoding the N-terminal part of the amino acid sequence shown in Figure 3 or 4, which N-terminal part terminates with a furin-recognition sequence.
 - 14. A host cell transformed with the recombinant vector of claim 13.
 - 15. A method for producing a soluble form of a membrane protein of interest having a C-terminal ectodomain, said soluble form essentially consisting of said ectodomain, which comprises:
 - obtaining nucleic acids encoding essentially the ectodomain;
 - fusing the nucleic acids in phase with the C-terminal end of the N-terminal part of the recombinant vector defined in claim 13;
 - having the fused nucleic acids to be expressed in a host cell in the presence of a culture medium, which host cell expresses or is made to express furin; and
- recovering said soluble form in the culture medium.
 - 16. A method as defined in claim 15, wherein said protein of interest is NL-3 or β -endorphin.
- 35 17. The soluble form of the metallopeptidase defined in any one of claims 1 to 4 which soluble form essentially consists of the ectodomain of said metallopeptidase.
 - A composition comprising the soluble metallopeptidase of claim 17.

- 19. An oligonucleotide derived from the nucleic acid defined in claim 4, which oligonucleotide has at least 12 nucleic acids in length.
- 20. An antibody directed against the metallopeptidase defined in any one of claims 1 to 4.
 - 21. A method for detecting the presence or amount of a metallopeptidase as defined in any one of claims 1 to 4 in a sample, which comprises the steps of contacting said sample with the antibody defined in claim 20, in conditions such that an immune complex is formed between said antibody and said metallopeptidase, and detecting the presence or amount of an immune complex as an indication of the presence or amount of said metallopeptidase in said sample.

22. A method of detecting the presence or amount of a metallopeptidase as defined any one of claims 1 to 4 in a sample, which comprises the steps of: contacting said sample susceptible to comprise a target nucleic acid with the nucleic acid defined in claim 5 or with an oligonucleotide as defined in claim 19 in conditions such that a hybridization complex can form between the target nucleic acid of the sample and said nucleic acid or oligonucleotide encoding said metallopeptidase, and detecting the formation of such a hybridization complex as an indication of the presence of said metallopeptidase in said sample.

	1 10 20 20									
NEP1-HU	= 20									
PEX-HUM	MEAFVGGTLVLG									
KELL-HU	MEGGDQSEEEPRERSQAGGMGTLWSQESTPEERLPVEGSRPWAVARRVLTAILIL.									
ECE1-HU	MSTYKRATLDEEDLVDSLSEGDAYPNGLQVNFHSPRSGQRCWAARTQVEKRLVVLVVLLA									
consens	M T P									
NEP1-HU	40 50 60 70 80 90 TIIAVTMIALYA.TYDDGICKSSDCIKSAARLIQNMDATTEPCTDFFKYACGGWLKR									
PEX-HUM	TILFLVSQGLLSLQAKQEYCLKPECIEAAAAILSKVNLSVDPCDNFFRFACDGWISN									
KELL-HU	.GLLLCFSVLLFYNFQNCGPRPCETSVCLDLRDHYLASGNTSVAPCTDFFSFACGRA									
ECE1-HU	AGLVACLAALGI.QYQTRSPSVCLSEACVSVTSSILSSMDPTVDPCHDFFSYACGGWIKA									
consens	L L C C L V PC DFF ACGGW									
NEP1-HU	100 110 120 130 140 150 NVIPETSSRYGNFDILRDELEVVLKDVLQEPKTEDIVAVQKAKALYRSCINESAIDSR									
PEX-HUM	* *** ** ** ** ** ** ** ** ** ** ** **									
KELL-HU	KETNNSFQELATKNKNRLRRILEVQ.NSWHPGSGEEKAFQFYNSCMDTLAIEAA									
ECE1-HU	** * * * * * *									
consens	NPVPDGHSRWGTFSNLWEHNQAIIKHLLENS.TA.SVSEAERKAQVYYRACMNETRIEEL N P G F L LK LE A KA V SCHOLE ALE									
	N P G F L LK LE A KA Y SCMNE AIE									
NEP1-HU	160 170 180 190 200 GGEPLLKLLPDI.YGWPVATENWEQKYGAS.WTAEKAIAQLNSKYGKKVLINLFVGTD									
PEX-HUM	DAKPLLHILRHSPFRWPVLESNIGPEGVWSERKFSLLQTLATFRGQYSNSVFIRLYVSPD									
KELL-HU	GTGPLRQVIEELGGWRISGKWTSLNFNRTLRLLMSQYGHFPFFRAYLGPH									
ECE1-HU	RAKPLMELIERLGGWNITGPWAKDNFQDTLQVVTAHYRTSPFFSVYVSAD									
consens	PL GWF TL Y F YV D									
NEP1-HU	220 230 240 250 260 DKNSVNHVIHIDQPRLGLPSR. DYYECTGIYKEACTAYVDFMISVARLIRQEERLPI.DE									
PEX-HUM	DKASNEHILKLDQATLSLAVREDYLDNSTEAKSYRDALYKFMVDTAVLLGA.NS									
KELL-HU	PASPHTPVIQIDQPEFDVPLKQDQEQKI.YAQIFRE.YLTYLNQLGTLLGG.DP									
ECE1-HU	SKNSNSNVIQVDQSGLGLPSRDYYLNKTENEKVLTG.YLNYMVQLGKLLGGGDE									
consens										
	T==1									

NEP1-HU	270 280 290 300 310 320 NQLALEMNKVMELEKEIANATAKPEDRNDPMLLYNKMTLAQIQNNFSLEINGKPFSWLNF
PEX-HUM	* **
KELL-HU	SKVQEHSSLSISITSRLFQFLRPLEQRRAQGKLFQMVTIDQLKEMAPAIDWLSC
ECE1-HU	EAIRPQMQQILDFETALANITIPQEKRRDEELIYHKVTAAELQTLAPAINWLPF
consens	M E A PER KT L P WL
	330 340 350 360 370
NEP1-HU	TNEIMSTVNISITNEEDVVVYAPEYLTKLKPILTKYSARDLQNLMSWRFIMDLVSS
PEX-HUM	* * * * * * * * * * * * * * * * * * *
KELL-HU	LQATFTPMSLSPSQSLVVHDVEYLKNMSQLVEEMLLKQRDFLQSHMILGLVVTLSPA
ECE1-HU	LNTIFYPVEINESEPIVVYDKEYLEQISTLINTTDRCLLNNYMIWNLVRKTSSF
consens	V L LNMW V
NEP1-HU	390 400 410 420 430
	LSRTYKESRNAFRKALYGTT.SETATWRRCANYVNGNMENAVGRLYVEAAFAGESK
PEX-HUM	LSRRFQYRWLEFSRVIQGTT.TLLPQWDKCVNFIESALPYVVGKMFVDVYFQEDKK * **
KELL-HU	LDSQFQEARRKLSQKLRELTEQPPMPARPRWMKCVEETGTFFEPTLAALFVREAFGPSTR
ECE1-HU	LDQRFQDADEKFMEVMYGTKKTCLPRWKFCVSDTENNLGFALGPMFVKATFAEDSK
consens	L FQ F GT PW CV G FV F K
NEP1-HU	440 450 460 470 480 490 HVVEDLIAQIREVFIQTLD.DLTWMDAETKKRAEEKALAIKERIGYPDDIVSNDNKLNNE
PEX-HUM	EMMEELVEGVRWAFIDMLEKENEWMDAGTKRKAKEKARAVLAKVGYPE.FIMNDTHVNED
KELL-HU	SAAMKLFTAIRDALITRLR.NLPWMNEETONMAQDKVAQLQVEMGASE.WALKPELARQE
ECE1-HU	SIATEIILEIKKAFEESLS.TLKWMDEETRKSAKEKADAIYNMIGYPN.FIMDPKELDKV
consens	L IR AFI L L WMD ET A EKA A GYP
	500 510 520 530 540 550
NEP1-HU	YLELNYKEDEYFENIIQNLKFSQSKQLKKLREKVDKDEWISGAAVVNAFYSSGRNQIVFP
PEX-HUM	LKAI KFSEADYFGNVLQTRKYLAQSDFFWLRKAVPKTEWFTNPTTVNAFYSASTNQIRFP
KELL-HU	YND.IQLGSSFLQSVLSCVRSLRARIVQSFLQPHPQHRWKVSPWDVNAYYSVSDHVVVFP
ECE1-HU	FNDYTAVPDLYFENAMRFFNFSWRVTADQLRKAPNRDQWSMTPPMVNAYYSPTKNEIVFP
consens	YF N LR W P VNA YS N IVFP
	= $=$ $=$ $=$ $=$ $=$ $=$ $=$ $=$ $=$

NEP1-HU	TI TO NOO SHOULD BE THE FOUND ON THE THEF DONGRN FNK DGDLVDWWTOOSASN F
PEX-HUM	*** *** * * * * * * * * * * * * * * * *
KELL-HU	AGLLOPPFFHPGY.PRAVNFGAAGSIMAHELLHIFYQLLLPGGCLACDNHAL
ECE1-HU	
consens	——————————————————————————————————————
NEP1-HU	620 630 640 650 660 670 KEQSQCMVYQYGNFSWDLAGGQHLNGINTLGENIADNGGLGQAYRAYQNYIKKNG.EE
PEX-HUM	KEKTKCMINQYSNYYWK. KAGLNVKGKRTLGENIADNGGLREAFRAYRKWINDRRQGLEE
KELL-HU	QEAHLCLKRHYAAFPLPSRTSFNDSLTFLENAADVGGLAIALQAYSKRLLRHH.GE
ECE1-HU	KRQTECMVEQYSNYSVNG.EPVNGRHTLGENIADNGGLKAAYRAYQNWVKKNG.AE
consens	KE CM QY N NG TLGENIADNGGL A RAY G E
	680 690 700 710 720 730
NEP1-HU	KLLPGLDLNHKQLFFLNFAQVWCGTYRPEYAVNSIKTDVHSPGNFRIIGTLQNSAEFSEA
PEX-HUM	PLLPGITFTNNQLFFLSYAHVRCNSYRPEAAREQVQIGAHSPPQFRVNGAISNFEEFQKA
KELL-HU	TVLPSLDLSPQQIFFRSYAQVMCRKPSPQDSHDTHSPPHLRVHGPLSSTPAFARY
ECE1-HU	HSLPTLGLTNNQLFFLGFAQVWCSVRTPESSHEGLITDPHSPSRFRVIGSLSNSKEFSEH
consens	LP L L QLFFL AQV C PE D HSP FRV G LSN EF
	740 750
NEP1-HU	740 750 FHCRKNSYMNPEKK.CRVW
PEX-HUM	FNCPPNSTMNRGMDSCRLW
KELL-HU	FRCARGALINPSSR.CQLW
ECE1-HU	FRCPPGSPMNPPHK.CEVW
consens	FC SMMP C W

PRIMER	SEQUENCE
(IA)	5'-TGGATGGAT/CGA/CIGG/AIACIA/CA-3'
(1B)	5'-TGGATGGAT/CGA/CIGG/AIACIA/CG-3'
(2A)	5'-A/GTIGTITTT/CCCIGCIGGIA/GT/AIC/TTA/TCA-3'
(2B)	5'-A/GTIGTITTT/CCCIGCIGGIA/GT/AIC/ITG/CCA-3'
(3)	5'-AIICCICCIA/TC/TA/GTCIGCIG/AC/TA/GTTT/CTC-3'
(4)	5'-GAT/CAAT/CTIGAT/CGAA/GT/CTIAAT/CTGGATGG-3'
(5)	5'-T/CT/CACCAIATICT/GA/GCATCG/TT/CTTCATIGGG/ATG-3'

TIEST 7

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ctgatatactaaccagagaaccaactagacaactatgagacatccaactatagattlaaggacttgactgactgactgagagcaccagggtccacttggggcacagattacagcattgagaacag agaccaggacagtgcaccagcttcagtgtgtactaggcatccagatccagctgcctctccctagcctggccttagcggtgtgtgccttccaccagaaccggttgagagaagtctgagagaccagtggg 30 ala GCC 60 val GTT 90 **a**3n AAC 367 120 100 150 300 180 Lys Aga AAC 240 asn thr ACG glu SP. AGC asn pro TTA leu len gln 367 T T leu CTG arg leu CTC ACC thr arg val GIG thr val asp GAC leu TTG glu GAG 11e ATC glu GAG hds CAT ser AGC ala GCT 11e ATA met ATG leu CTG asp arg ST ST gln leu leu TTG tyr phe leu leu leu] trp វិតីព ala GCC 11e ATC leu ser TCC CTG gln glu GAG AIC AIC ser TCC ala thr ACT val GTG pro arg CGA GIC Val leu CTG phe ala gg 14 S ser TCC glu GAG tyr glu his CAT his CAC val GTG 11e ATA asp GAT ser TCT leu arg ala ပ္ပ met ATG leu val GTG arg glu GAG asp GAC glu Gaa AGC AAA 36Z leu CTG leu CTG leu CTG cys TGT len CTG arg AGA trp 166 JGG arg CGG 91y 666 3er AGC ser AGC trp TGG val GTG lys AAG lys AAG asn AAC val GTA tyr Tat thr ACT pro CCA 91y 660 glu GAG 91y 666 leu CIC gln 1ys AAG glu Gag len TTA thr Acc 91. 88. 173 848 110 ATC 91y GGC asp GAC his Cac leu val GTG leu CTC thr cys TGC STO val GIG AAC AAC met asp GAC phe Pro ala GCC C.Y.s 160 11e ATC ser AGT thr Acc asn 91y 660 leu CTG 116 ATC tyr a Bp GAC val GTT gln glu GAG trp 166 or S gg asp GAC gln glu GAG asn AAC asn AAC glu GAG 11e ATC Ser TCC 1ys AAG ser leu CTG tyr TAC met ATG trp IGG phe gla lys AAG 917 GGG lys Aa phe glu Gag 1ys AAG cys TGC 1eu C70 phe 1ys AAG 11e ATA leu CTG aan AG asp GAC asp TG. 361 asp GAC tyr Tar arg CGG ser AGC ser glu Gaa arg arg met ATG 11e ATC tyr Cys TGT tyr TAC ser TCA cys leu CIG leu CTC tyr TAT ala GCC glu GAG trp 196 phe GAT pro CCC 11e ATC 1eu CTA val GTG val arg CGG val GTC 91.y 660 arg Agg AAC asp GAC g th pro arg CGG 3er TCC ala GCA leu CTC gly GGT arg Agg phe lys AAG trp 199 pro CCA arg Agg glu arg GAG AGA leu CTG ala GCC ser TCG val ala 917 GGT AAC met ATG thr Act arg gg gg ser AGC 1ys AAG phe 91.y 668 915 val val GTG 133 843 asp GAC tyr glu GAG val GTA 92 88 leu TTG met ATG 11e ATA val GTA met ATG arg CGA val GTG met ATG Ber TCG thr gg 5 181 271 541 361 151 631 721

from mouse CDNA ML-1 e O Saquence

300 CAT 330 330 390 420 420 420 420 420 11ys AAG 510 510 thr arg val leu glu asp GAC Ber asp GaT 1ys AAG val GTC glu GAG glu val GTG arg glu GAG 1ys AAG gha leu leu 11e ATC leu CTG age See glu trp Igg val GTG Ber glu GAG his Cac ser AGC gln glu leu glu val GTA thr phe asp Lys Aaa arg AGG asn AAC leu glu GAG asp GAT leu CTG thr ala GCC met asn AAT gln arg AGA val GTG met ATG Pro tyr 917 arg trp Igg asn ala GCC asn glu GAG leu CTG phe asn tyr 1ys AAG asn AAC asp GAT asn AAT ដូសូ ala GCG asp GAC leu CTG 95 95 95 leu CTG 11e ATC leu CTG glu asn AAC met ATG met ATG glu GAG met ATG ala GCG tyr glu GAG leu TTG lya Aag tyr glu GAG arg val GTC thr 1ys AAG leu CTC asp GAT 11e ATT leu CIC phe glu GAG his Cac 95 83 arg CGG arg ser TCC leu tyr TAC asn ala Ge arg tyr val GTG ala GCA tyr Tac 91.y 660 ASD asp GAC gln asn AAT val GTG leu Ser ser TCA asp GAC val asp Gat pro leu val met ATG ala GCC ser tyr val GTG ala GCC val tyr TAC gly gga val GTG **ala** GCC thr leu TTG ser AGC arg Ber phe TTT 917 GGC agn ala GCA 30r val Grc val GTG asp ala GCG glu val GTG 11e ATT glu GAG ala GCT glu asp asn AAC 11e ATT glu GAG met ATG ser TCC gln phe TTT 91*y* 666 Lys AAG his Car gh 11e ATC Lys Aaa asn AAC arg gyn tyr 11e ATC ser TCC arg Agg 11e ATA asp phe Ber 11e ATA arg leu 118 ATC 1eu CTG 1.78 8.88 phe glu GAG arg asn AAC lys AAG 11e ATA asp trp Igg AAC glu GAG 1eu CTC leu CTG glh glu GAG val GTC asn glu GAG leu gg. 42 thr Asn ser Agc tyr TAT 11e ATT net ATG tyr TAT agn asp GAC ign ign glu leu CTG ser AGC leu ala GCC phe gla 173 848 val GTC asn AAC leu CTG aer AGC val GTC glu Lys AAG asp GAC thr arg AGG thr phe tyr 917 660 cys TGT arg Aga gla leu TTG val GTG leu ala GCC 917 Pro CCC 11e ATT gyn gla Gag val GTC Ser lys AAG met ATG asa AAC 1ys AAG 11e ATC arg CGA arg thr ala GCC gra ser TCC thr ala GCC leu CTG 917 asp GAT ign ign ser AGC 1ys AAG tyr TAC arg CGG ala GCC leu CTG gly ggt leu CTA tyr lys AAG arg lys AAG gra leu 811 901 166 1081 1111 1261 1351 1531 1621

F=== 3 (cont'd)

val

ala GCC

phe

gla

pro

gln CAG 750 Cys

hts Cac

phe

ala GCA

glu Gag

Ser

leu CTC 690 leu CTG 720 his CAC his CAC thr arg CGA 91y 666 arg CGG Ser gln 11e ATT ala GCC phe asp val GTG Ser TCG 917 968 lys Aaa ATG phe asn 91y 660 91y 666 asn AAC val 91y 660 ile ATC Ser asn asp GAT 91y 660 trp gln CAG ala GCT 91y GGG **trp 1**GG asn AAC leu phe asp GAC trp TGG asp ser leu asn proposer leu CTG ala GCA arg CGG met ATG leu leu CTA asn AAC glu GAA tyr gln 91y 660 trp 166 ala GCT or S asn AAC ser lys Aag gla GA 1ys phe TTC tyr asp GAC asp GAC asn AAC ala GCA lys AAG phe gly GGC gla RGC asn AAC tyr arg phe gla gly val arg phe gly GGT tyr TAT 9 CC CC CC asn AAT 11e ATC 91y 66a asp Pro CCG met ATG asn AAC gln CAG asp asp cys TGC ဌဌ leu phe TTT gg gg ala GCC 11e ATT 91.7 93.0 93.0 ser TCG 11e ATT his CAC gln asn ala GCA g th gln CAG glu GAG er S 11e ATC g CAA 91y GGG 1711 1801 1891

arg phe TTC tyr TAT 91.y 660 ser TCC ರ್ಗ ೧೭೩ 917 666 leu CTG asn AAC Cys TGT gln trp IGG val GTG leu CTA gln ser TCA ala GCC 91y 660 tyr TAT leu CTG asn val GTG 11e ATC arg phe tyr phe lys AAG leu leu gln pro ala GCC ser AGT tyr TAT his CAC th Sc val GTC Leu asp GAC asn AAC thr len CTG lys AAG 11e ATC gly gga ser TCC Pro

P. CAC gly ser GGC AGC arg CGA pro

TAG CCAAGGCTGAGCTATGCTGCGGCCCACGCCCCGCCACCCCAGAGGCTTCGCGAATG 136 tr 11e ATC Cys arg TGT CGC arg pro met lys CCC ATG AAG met ATG Pro

2473

TTC

2925

GIGIAGCIGGCAGAAIGIGCAGGICITIGCCIGAAGGCCACCGGAGCCAGCCAGCCCICCGCGCCCAAGCCTAAAGIGIAGCCACCCGCCCACACCGGGAIGAGIGGGGCGGGT

_ **3** (cont'd)

2161

2071

humans

from

CDNA

NL-2

of

Sequence

8/22

glu GAG 88 CTG GTG ACC ala GCA 118 his CAC ser TCG 178 gln CAG 208 glu GAG his CAC 268 tyr TAC 298 7 91y leu leu leu leu leu leu leu val GGG CTG CTG CTG CTG CTG CTG GTG ; gln ile ATA arg asn ser leu CTG arg CGG ည္ဟ val leu TTA val GTG arg glu GAG glu GAG 91y GGC glu ser met ATG phe cys TGC leu leu CTG arg trp TGG ser arg CGG asp $\frac{cys}{16c}$ 91y 660 trp val GTG lys AAG glu GAG asn val GTG glu GAG leu CTG pro 91y GGC ala GCG glu GAG leu CTC gln lys AAG gln arg thr gly GGA lys AAA ile ATA gly GGA asp arg CGG val GTG ser thr Cys TGC leu val GTG val GTA asp asn leu ala GCT cys TGC ala GCA ile ser thr asn ser Cys TGC leu val GTC phe TTT val GTC gln CAG glu GAG trp TGG gly GGC ser ogly phe leu glu gly GGG rrc crg GAG GGG c arg glu GAG gln CAG glu GAG ASD asn ile gly GGC asp pro Ser tyr leu CTG met ATG trp **T**GG phe TTC asn arg leu val phe glu GAG cys TGC arg AGG leu CTC phe pro gln glu GAG asp asp ser TCC asp asp tyr leu lys AAG gla asp GAC arg arg me t ATG ile tyr asn AAC arg ala gly gln lys arg pro GGT GCA GGG CAG AAG CGC CCG (\mathfrak{g}_{1} y GGG ala GCC cys TGT leu tyr ala GCG leu glu GAG ala GCA arg glu GAG pro CCG val GTC leu CTG val GTG val GTC arg asp GAT ATG GTG GĀG AGC GCC GĞC CGT GCA GĞG CĂG AAĞ arg pro glu GAA asp thr pro glu GAG arg ser asp ile ATC thr phe arg trp arg pro CCC arg 917 GGG ala GCC thr ile ala GCC \mathfrak{g}_{1Y} ASD met ATG leu CTG tyr arg pro Ser lys AAG gly Gga phe TTC leu TTG 91y GGC leu pro asp GAC tyr glu GAG val GTG gln leu TTG thr gly val GTC lys Aaa met ATG arg val GTG val ser thr ala GCC qlu ser ala gly GGT arg asn ser glu GAG ala GCT asn pro val GTG leu TTG lys AAA gln asn pro leu TTG met ATG gln ser ala GCC val GTA leu thr arg ile ATC leu asp val GTG met val val GTG phe ile ATC glu asp asp ala GCG ile ATA met ATG leu CTG thr arg pro lys AAG leu tyr leu CTG phe 999 ala GCC arg ala GCC ile ATC ala GCC leu CTG gln ile gln GTG ala GCT glu GAG ala GCC val GTG thr pro arg CGG ile ATC leu CTG 91 181 271 361 451 541 811 631 721

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. CTG 358 1eu CTG 388 388 asn AAC 418 phe TTT 448 arg AGG 91Y GGC 508 91u GAG 538 91Y GGC 568 Pro 91y GGA leu gln leu CTG val GTC leu glu val SOL met ATG lys AAG ile ATA ala GCG tyr glu leu CTG Lys AAG tyr 91y 664 arg CGG ile thr lys AAG leu asp ile ATC leu phe his 1ys AAA arg arg Ser leu Tyr BC asn ala GCG asn AAC tyr tyr val GTC ala GCC 91y GGC thr asp GAC leu TTG gln AAT leu TTG ser ser asn val GTG glu GAG pro leu CTG ala GCC val Grc ala GCC ser TCC tyr TAC val GTG ala GCC val GTG his Ser val ag g ile ATC leu thr arg asn phe 917 asn ala SSS Pro val GTG val asp thr glu GAG val GTG ile ATC glu ala GCG gla CAG asp thr ile ATC asp met ATG thr g Gac phe TIT 917 666 glu his gln ile lys AAG asn arg glu Geg lys AAG ty: ile ATC arg ile ATA asn phe Ser val GTG arg leu CTG ile ATC ser AGC glu GAG phe glu arg asn lys AAG ile ATC asp GAC trp igg phe glu GAG leu leu gln val GTC asp ser AGC glu leu CTC phe gln thr asn Ser tyr TAC ile ATT met ATG asn 38r TCA Pro pro trp TGG gln leu 91y 660 leu ala GCC phe er Services Pro val asn leu CTG ser val GTG glu Gaa 1ys AAG asn AAC gln asp GAC thr tyr TAC phe TTT gly GGT cys arg glu leu CTG leu CTC val ala GCC 91y 66a Pro ile ATT glu GAA val GTC gln asn lys AAG ile lys AAG 1ys AAG ile arg arg met ATG ala GCG Ser glu Gas 91y 666 ala GCC leu gly GGC asp $\operatorname{trp}_{\operatorname{TGG}}$ ser tyr Tac lys AAG arg ala leu 91y GGC tyr TAT leu CTG arg lys AAG lys AAG glu leu CTT pro gln phe val GTG val GTG ser val lys AAG glu lys AAG phe thr gln val leu glu GAG asp Ser asp GAC arg AGG val GTA glu GAG glu GAG val GTG arg gly GGA glu leu Cic leu ile Arr leu CTG gln glu GAA trp TGG val GTG pro glu gla arg CCC 3er AGC glu GAG leu glu GAG val met ATG phe asp GAC arg AGG arg asn leu glu GAG asp leu thr ala GCG met ATG asn AAC g Ge arg CGA val GTT glu GAG pro tyr 91.y GGC glu GAG met ATG ala Scc asn स्म गुद्ध 901 1081 1171 1261 991 1531 1621 \Box

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Ser	658 asn AAC	688 glu GAG	718 tyr TAC	748 ala GCC	GTG	၁၁၅	GGA	AAG	
ig ig	gln	ala GCA	ser Toc	ala GCC	GCT				
tr 166	glu Gaa	met ATG	91y GGG	leu CTG	282				
asp Gat	asp GAC	trp TGG	cys TGC	asn	TGC				
met AIG	ala Q	1ys AAG	trp 166	gln					
met AIG	leu CTG	leu CTC	val GTG	leu CTG	AGG	သည		TCC	
asn	asp GAC	tyr TAC	gln CAG	ser	₹ CC			GTG	
91y GGC	मु है	ala GCC	ala GCC	gly GGG	ter TAG	CGT	gcy	ACT	
asn	ser	lys AAG	tyr TAT	leu CTG		GTA	TGA	TCA	AAA
lys AAG	なる	tyr TAT	asn AAC	val GTA	val GTG	AGT	GGA	CAA	AAA
asp GAC	asn	ala GCC	ile ATC	arg AGG	arg	သည	CAG	AGA	AGA
phe	91y 660	gln	phe TTC	tyr TAC	cys TGC	CGA	ACC	TAG	CAA
asn	tyr Tac	arg CGG	phe	lys AAG	arg CGA	SSS	GAC	CTG	TTT
arg CGG	gln	val GTG	leu	leu CTG	glu GAG	TAG	CCI	ACT	IGC
91 <u>y</u> 660	tyr	91 <u>y</u> GGG	gln	pro	1ys AAG	AGC	CTG	ACA	AAA
asn	ile ATC	917 663	glu GAG	ser	P.70	TGC	CAC	GIC	AGT
asp GAC	met ATG	asn	his CAT	his CAC	his	AGG	AGC	SCC	CTA
asp GAC	973 760	asp GAC	thr	val GTC	met ATG	CGA	T GC	၁၁၅	GAG
phe	glu	ala GCT	leu	asp GAC	Pro	GTG	999	AGT	CTT
917	3er TCA	ALT	asp GAT	thr	thr	TCT	CTA	CIC	gya
his Go	gla	AAC	leu CTG	lys AAG	gly GGC	GCA	299	999	TCA
ACG	glu GAG	glu Gaa	91y GGC	ile	arg	GAG	CCT	909	GCT
AIC	arg 000	91y 666	pro CCC	ser TCC	ala GCC	TCG	909	ပ္ပ	ACA
gly met val ile gly his glu GGG ATG GTG ATC GGG CAC GAG	phe TIC	leu CTT	leu	gln	cys TGT	IGC	GCC TCT GCG	ACA	TCT
# 5 -	gln his CAG CAC	asn thr AAC ACC	gln	ile	his	၁၅၁	၁၁၅	TCA CAG	GTA
498	of contract of the contract of	AAC	gln	ala	phe TTC	CCC ACC	CAG	TCA	CCA
i i	thr Acc	phe	asp GAC	phe	thr	သသ	GCC TGC	CCI	CIT
7 C	B BBE	gly GG	1ys	glu	asp GAC	CCC ACG		CCC	TGT
≥ S	phe TTC	asn AAC	gly : GGC	pro CCC	ala GCA		CCT	GAG	CAT
	asn 1 AAC	val I GTG	gly I GGT	arg CGG	phe	555	AAG	ဗ္ဗ	GTG
1801	1891	1981	2071	2161	2251	2341	2431	2521	2611

T== 4 (cont'd)

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Sequence of NL-3 cDNA from human

11/22 ცვ 61C arg CGC 52 leu CTG 112 112 142 GAC GAC pro CCT 202 1eu CTA 232 1eu CTG 262 arg CGC val GTG 322 asn AAC 990 CAG Ser 91y GGG ala GCC phe Pro 91y 666 met leu ile arg ala GCC GCT ည္ဟ val ser TCC leu ile arg gly GGT pro arg CGG val tyr leu ပ္ပည္သ GTG arg met ala GCT ala GCC 91y 666 asn arg tyr ala GCA gln ၅၁၁ CAG ala GCC ala GCT his ala GCC pro pro leu arg ala GCA gln TCA ပ္ပပ္ပ val GTC 91y 666 ala GCG arg arg arg gly GGC asp ser leu CTG glu GAG SSS GIA glu GAG thr leu ala GCG arg CGG ala GCG leu trp ser val val GTG GCT ပ္ပပ္ပ gha ala GCC ile ATT phe TTC leu leu arg arg asn gln SSS CAT phe ala GCC ser ala GCC trp TGG leu glu GAG ala GCG arg glu leu CTG CGA ပ္ပပ္ပ glu GAG arg cys TGC lys AAG gly arg ile ala GCG asp ile ATC Ser SCC CAC asp GAT ala GCG leu CIC arg 91y GGC arg glu GAG val GTC asp glu GAG TCG ပ္ပဗ္ဗ tyr TAC ala GCT 91y GGC glu GAG cys TGC leu CTA arg 91y GGG leu glu GAG gln ပ္ပဗ္ဗ S his CAC 91y GGC ala GCC pro ala GCC met ATG arg pro ser asp ala GCC GAG GAG GCC ပ္ပ ala GCG leu TTG ala GCC cys TGC phe glu GAG asp arg val gln lys AAG ပ္ပမ္မ Pro CCG thr phe **91** y GGC ser TCG glu GAG glu leu thr ala GCT gln TCA leu CTG phe val GTG glu GAG tyr asn Cys TGC glu GAA glu GAG leu leu CAC 299 ser TCG leu CTG 91y 660 pro phe gg Gg ser ala GCG ser TCG tyr val GTG ပ္ပ 733 tyr TAT Pro 91y GGG cys TGT asp glu GAG arg gly GGC phe leu CTG ala GCT GAG ည္ပည Pro Pro ser ala GCC gln 91y GGC phe thr leu CTC asp 91y 660 CIG 929 pro leu leu 917 Cys ile phe ala GCG leu arg ala GCA ອອິ CAT glu GAG Ser leu CTG 91y GGC Pro ala GCA ala GCC glu asp ala GCC gly GGT GCT ည္ဟ met ATG ala GCC cys TGC 91y 66c asp ala GCG arg trp 166 ala GCC pro leu CAG ပ္ပ ည္ပ val 91y 666 ala GCC ile ATC ile val GTG Ser leu leu 91y 660 ACG ပ္ပ ပ္ပပ္ပ arg glu GAG ala GCG Ser thr lys 91y GGG tyr TAC thr Ser 990 CGG 700 arg CGC ala GCG ala GCC val GTC 91y 66c arg cys TGC val GTG leu leu CAC CTG ပ္ပ gly arg CGG pro asp tyr TAT gln asp 91y 660 91y 666 val AGA ეცე ცვ 91y 666 asn 91y GGC leu CTG thr ala GCC glu gln asp 750 ეც ala GCG trp 166 leu asn leu ala GCG ile ala GCG gln glu GAG ACC GCT ACC ala GCC arg tyr 91y 660 lys AAG 91y GGC val GTC lys AAG asp ပ္ပမ္မ ပ္ပပ္ပ 7357 Pro lys ala GCC cys TGC asp glu GAG tyr ile ATT 91y 660 phe 5 271 361 151 541 631 721 811 901 1081 991 000

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352	20	382	ATG	412	his	CAC	442	leu	TTG	7/5	gIn	S	205	lys	AAG	532	glu	GAG	562	thr	ACG	592	leu	CIG	622	gly	666 67,7	770
† 7	ACC	1	TAC		glu	GAA		cys	1 GC		gin	CAG		ala			his	CAT		30r	100		Ë	ACC		Ę,		
	ATC	ر د	GAR		ser	AGT		val	GIC	•	val	GIG			990			GIC		lys			pro	ပ္ပ		asp	S S	
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<u>ر</u> د	CAG	<u> </u>	909		val	GIC		ala	ದ್ದಿ		aTa	ပ္ပ			GCH			TII		val				CIG		Q.		
1	CTG	<u> </u>	CTG		val			leu	CIG		τys	AA		ala				GAG		glu	GAG			ATC		gly		
5	CAG	[CTG		va]	GIG		glu	GAG		ser	AGC		arg			tγr			gln			3 12			hia		
=	944	(eA	GTG		val	GIG		gln	CAG		aTa	ပ္ပ		돢			glu				ပ္ပံ		ala			th.		
,	CIG	(e)	GTG		arg	ပ္ပ		pro	CCA		aTa	GCI		gJn	S. C.		lγs			118			pro			leu		
۲ د د	ACG		GAG		trp			lys	AAG		ser	17.	,	ala glu	ပ္ပ		asp				AG		phe			glu	3	
	GTG	2 2	GAG		Val	GTG		asb	GAC	-	bue	TIC		asb	S		val			ΙΫ́з		•	Val	GIG		his		
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נו נו	AAC		GAG		tyr			g1y	ပ္ပပ္ပ	•	ara	S S		Į,	TGG CG		asb			361			gTn		!	110	ALL	
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đ t	ATG	phe	TTC		his			met	ATG		אפא	<u> </u>		Ten	CHO	,	lys	¥		gln	S		₹₹	AG A	:	th F	ACC	
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	AGC		GAG		Val			gIn	CAG	-	T e T	CI CI		gIu	S		Ten				ပ္ပ		ord or			110 110	YIV.	
	GTC		CAG		arg			ala	GCA	-	7	ပ္ထ		leu	CIG		phe			phe			Ten	E E		775 775		
	GAT		TTC		his			leu	CIG	-	đΤδ	ក		gly gin arg	ပ္ပ		dee.			arg			F.			91Y		
π π	KSD :		ATC		pro			glu	GAG	,	naT	C C	,	<u>п</u> Тб	5		pro	ပ္ပ		110			ζ			tyr 3 Yr	141	
	990		CAG		thr			his		-	לי ד	ပ္ပ		δTδ	ပ္ပဗ္ဗ		tyr			30r			414			מפש		
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20.	9.6	leu	CTG		ile	ATC		glu	GAG.	1	bud	S T		tyr	TAC			AIG	:	118	ATC	•	bro din ara	ပ္ပ	,	phe pro gin	3	
+	TAT	Tr.	TGG		l leu	5		arg		لـ	nıs	ប្រ		Τλs	AAG	•	met val	ត្តវិត		lys asn	Z	•	5	ទ		Ord C	5	
-	GTG TCA GAG	108	AAG		gln	S		bhe	TTC		arg	ខ្ល	:	Te	ATC		明日	AIG	,	Įχg	Z		Д. О	ខ្ល		pha) [
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	1 ATC	his	CAC		gln			len	r L	•	91.	1531 GGC CAG GCC AAT CGC CAC TIT GGC ATG		len	CTA		leu			1Ys				16G		Ş.		
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t z = 5 (cont'd)

914 914 by the target ser gly and lead his trip this glu ala ser tyr ser arg phe lead arg lys ala glu cys ile val argon cor card cord and cord cord cord cord cord cord cord cor					/3/	12	2			
94.9 gin tyr asp arg ser gly asn len his trp thr glu ala ser tyr ser arg phe leu arg lys ala glu cys ile val ecc cor crox coc crox and crox crox crox crox crox crox crox crox	arg CGT	682 1eu CTG	712 Phe TTC	742 tyr TAC				SAG	SCT	
94.9 9th tyr asp arg as gly asn leu has trp thr glu ala ser tyr ser arg phe leu arg lys ala glu cys ile GCC CRC for AGC CRC for CRC AGC CRC TRC AGC CRC TRC AGC CRC AGG CRC TRC AGC CRC AGG CRC AGG CRC AGG CRC AGG CRC CRC AGG CRC AGG CRC AGG CRC AGG CRC AGG CRC CRC AGG CRC CRC CRC AGG CRC CRC CRC AGG CRC CRC CRC CRC CRC CRC CRC CRC CRC C	val									
GCC CAG TAT GAC CCC TCA GGG AAC CTG CTG AGG GGC TCC TAC AGG CGC TTC TAC AGG CCT TTC CTG CGA AAG GCT GGG TGG TGG CAG CAG TAT GGC CTG TAC AGG CTG CTG CGG AAC AT CTG CTG AAG GCT GGG GGG AAC AT CTG CTG AAG GCT ATG GGG GGG AAC AT TTG GCT TAC TAC TAC TAC TAC TAC TAC TAC TAC T	ile Arc	leu CIC	leu CIC							
GCC CAG TAT GAC CC TCA GGG AAC CTG CAC CAG TGG AGG GGC TCC TAC AGG CCT TC CTG CCA AAG GCT CAG AGG GCC TCT TAC AGG CAC CTG CAG CAG CAG GAG GCC TCT TAC AGG CAG AAC CTG CAG CAG GAG GCC TCT TAC AGG CAG AAC CTG CAG CAG AAC CAG CTT GGG GAG AAC ATG GCA CAT TAC AAC TTG CAG TAC AAC CAG GTG AAC GCG TT GGG GAG AAC ATG GCA CAT TAC GCA CAT TAC AAC TTG TAC AAC TTG TAC AAC CGG GAG AAC CAG CTT GGG GAG AAC ATG GCA CAT TAC GCA CAT TAC AAC TTG TAC AAC TTC TAC TTG TAC AAC TTC TAC TA	cys TGC	91y 660	g g							
GC CAG TAY GAG ALG CAC TCA GCG AAC CTC CAC TGG TGG AGG GGC TCC TAC AGC CGC TTC CTG CGA AAG GCT leu tyr asp asn phe thr val tyr asn gln arg val asn gly lys his thr leu gly glu asn ile ala asp met CTC TAT GAC AAC TTC ACT CTG CAG GTG AAC GGG AAA CAC GTT GGG GAG AAC ATC GCA GAT ATG AAC TTC ACT TAC AAC TTC AAC GTG GAG AAC GTG AAC GGG AAA CAC GTT GGG GAG AAC ATC GCA GAT ATG AAC TTC CAC GTG TAC GGG GAG AAC GTG GAG GAG CAC GAG GAG CTC CAG GTG GAG AAC ATC GAG GAG AAC ATC GAG GAG AAC ATC GAG AAC ATC GAG GAG GAG CAC GAG GAG CAC GAG GAG CAC GAG GAG	glu GAG	91y 660	asp Cac							
GC CAG TAT GAC CGC TCA GGG AAC CTG CTG CTG CTG CTG CAG GAC GAC GTG TTG TAC AGC CGC TTG CTG CTA AAG CTG CTG CTG CTG CTG CTG CTG CTG CTG CT	ala GCT	me t ATG								
GCC CAG TAT GCC TCA GGG AAC CTG CAC TGG TGG AGG GCC TCC TAC AGC CGC TTG CTG CGA Lleu tyr asp arg ser gly asn leu leu his trp trp thr glu ala ser tyr ser arg pea leu asn gln arg val asn gly lys his thr leu gly glu asn ile ala cTG TAC TAC TAC TAC TAC TAC TAC GGG GTG AAC GGG AAA CAC TTG GGG AAC TTG GCG AAC TTG CAC TTG AAC GTG GTG GGG GTG AAC GGG AAA CAC TTG GGG GAG AAC TAC GCG TAC TAC GAG TAC GTG TAC GAG TAC GTG TAC GAG CAC GAG CAC GAG CAC GAG CAC GAG CAC TAT CAC GAG TAC GTG TAC GAG CAC GAG CAC GAG CAC GAG CAC GAG CAC GAG CAC TAC TAC GAG TAC	lys AAG	asp Gat		1ys Aag	me t					
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GCC CAG TAT GAC CGC TCA GGG AAC CTG CTG TGG TGG AGG GAC TCC TAC AGC CGC TTC leu tyr asp asn phe thr val tyr asn gln arg val asn gly lys his thr leu gly glu asn CTC TAT GAC AAC TTC ACT GTC TAC AAC CGG GTG AAA GGC AAA CAC ACG CTT GGG GAG AAC ala tyr his ala tyr gln lys trp val arg glu his gly pro glu his pro GTT GGG GAG AAC ala tyr his ala tyr gln lys trp val arg glu his gly pro glu his pro GTT GGG GAG GAG cC TAC CAC CAC TAT CAC AAC TGC GGC GAG CAC GCC CCA GAG CAC CTT CCC GGG CTC ile ala phe ala gln asn trp cys ile lys arg arg ser gln ser ile tyr leu gln val leu ATT GCC TTT GCC CAC AAC TGC TGC ATC AAC GGG CGG TCC CAC TCC ATC TCC TCC TGC TGC arg val leu gly ser val ser gln phe glu glu phe gly arg val leu his cys pro lys val AGG GTG CTG GCC AGT GTG TCC CAC TTT GAG GAG TTT GCC CGG GTT TTA CAC TGT CCA AAG GTC Ser val trp ter TCC GTG TGG CTG CCC CAC ACC TGC TGC TGC TCC CCA GGG CCC CCA GGA CCC CGG TTTA CAC TGT CCA AAG GTC GGT GCC GGC GGC CTG GGC TTT GAG GAG TTT GGC CCC CCA GGA CCC CGG TTTA CAC TGT CCA AAG GTC Ser val trp ter TCC GTG TGG CTG CTG CTG CTG TGC TGC TGC	leu CTG	ile	lys Aag		aer TCA					
GCC CAG TAT GAC CGC TCA GGG AAC CTG CAC TGG TGG AGG GAC GCC TCC TAC AGC CGC leu tyr asp asn phe thr val tyr asn gln arg val asn gly lys his thr leu gly glu CTC TAT GAC AAC TTC ACT GTC TAC AAC CGG GTG AAC GGG AAA CAC AGG CTT GGG GAG ala tyr his ala tyr gln lys try val arg glu his gly pro glu his pro leu pro arg GCC TAC CAC CTT CAC AAC TGG GGG GAG CAC GGG CAA CAC CTT CGC GGG ile ala phe ala gln asn try gln lys arg arg asg ger gln ser ile tyr leu gln val ATT GCC TTT GCC CAG AAC TGC TCC ATC AAG GGG GAG CCC CAC GAG CAC CTC CAC GTG GGG avg val leu gly ser val ser gln phe glu glu phe gly arg val leu his cys pro lys AGG GTG CTG GGC AGT GTC CAG TTT GAG GAG TTT GGC CCC CCC GTG TTA CAC TGT CCA AAG Ser val try ter TCC GTG TGG CCC GCT TGC GCC TGC ACG TGC CCC CCC CCG GTT TTA CAC TGT CCC CCT GGT GCC AGG CCC GCT TGG GGC TGC ACG TGC ACG GTT TTA CAC TGT CCC CCC CAG GTG GGC AGG TTT GGG GGG GTT CCA GGG GTT TTA CAC TGT CCC CCC CAG GTG TTG GGC CCC TGC ACG TTT GGG GAT TTT GGG GGT CCC CCA GAG TTT CCA GGA CCC CCC CCC CCC CCC CCC CCC CCC	phe	asn AAC	leu CTC							
GCC CAG TAT GAC CCC TCA GGG AAC CTG CTG TGG TGG AGG GAC TCC TAC AGG leu tyr asp asn phe thr val tyr asn gln arg val asn gly lys his thr leu gly CTC TAT GAC AAC TTC ACT GTC TAC AAC CGG GTG AAC GGG AAA CAC AGG CTT GGG ala tyr his ala tyr gln lys trp val arg glu his gly pro glu his pro leu pro GCC TAC CAC TAT CAC AAC TTC AAC GGG GTG AAC GGG CAA CAC CCA CTT GGG ala tyr his ala tyr gln lys trp val arg glu his gly pro glu his pro leu pro GCC TAT GAC AAC TAC AAC GG GGG GGG GGG CCA GAG CAC CCA CTT GCG ile ala phe ala gln asn trp cys ile lys arg arg ser gln ser ile tyr leu gln ATT GCC TTT GCC CAG AAC TGG TGC ATC AAC GGG CCG CCA GGG CCA ATC TAC CTG CAG arg val leu gly ser val ser gln phe glu glu phe gly arg val leu his cys pro AGG GTG GGG CAG GC CAG TTT GAG GAG TTT GGC CCG GTT TTA CAC TGT CCA AGG GTG CTG GGC ACT GTG TCC CAG TTT GAG GAG TTT GGC CCG GTT TTA CAC TGT CCA AGG GTG CTG GGC ACT GTG TCC CAG TTT GAG GAG TTT GGC CCG GTT TTA CAC TGT CCA AGG GTG GGG CCC GCT TGG CTG CCC TGC ACG CCT CCA GGA CCC CCA GAG CCC CAG GGT GAG CCT GGG CTG GCG GTG GTG GTG CCT CCA GGA CCC CCA GAG CCC CAG GGT GAG GCT GGG CTT TGG GGG GCT TTT GCT GCA GTG CCC CCA GAG ATC CCC CAG GGT GAG GCT GGG CTT TGG GGG GCT TTT GCT GCA ATT TAC TGG GGT CCC CAG ATT CCC CAG GGT GAG GCT GGG CTT CGC TGT CTT CTT GCT GCA ATT TAC TGG GGT CCC CAG ATT CCC CAG GGT GAG CCC CAC CTT CGC TGT CTT CTT GCT GCA ATT TAC TGG GTC CAG ATC TCC CAG GGG TTT GCT TTT GGT TTT CTT GCT TTT GTT G	arg ೧೧೧									
GGC CAG TAT GAC GCC TCA GGC AAC CTG CTG CTC TCG TGG ACG CAG GCC CAG TAT GAC CGC TCA GGC AAC CTG CTG CTG TCG TGG ACG CAG GCC CTC TAT GAC ACT TTAT ACT TTAT ACT TTAT ACT TTAT CAC TTC TAT TAT			pro CCC	gln	Pro Ch		TCA			ATC
GGC CAG TAT GAC GCC TCA GGC AAC CTG CTG CTC TCG TGG ACG CAG GCC CAG TAT GAC CGC TCA GGC AAC CTG CTG CTG TCG TGG ACG CAG GCC CTC TAT GAC ACT TTAT ACT TTAT ACT TTAT ACT TTAT CAC TTC TAT TAT					cys TGT		GZA			TAA
GGC CAG TAT GAC GCC TCA GGC AAC CTG CTG CTC TCG TGG ACG CAG GCC CAG TAT GAC CGC TCA GGC AAC CTG CTG CTG TCG TGG ACG CAG GCC CTC TAT GAC ACT TTAT ACT TTAT ACT TTAT ACT TTAT CAC TTC TAT TAT				TAC	his			ပ္သပ္ပ		AAA
GGC CAG TAT GAC CCC TCA GGG AAC CTG CAG CAG TGG TGG ACG CAG leu tyr asp asn phe thr val tyr asn gln arg val asn gly lys CTC TAT GAC AAC TTC ACT GTC TAC AAC CGG GTG AAC GGG AAA ala tyr his ala tyr gln lys trp val arg glu his gly pro glu GCC TAC CAC CAC TAT CAG AAG TGG GTG CGG GTG CAC GGG CAG GCC ile ala phe ala gln asn trp cys ile lys arg arg ser gln ser ATT GCC TTT GCC CAG AAC TGG TGC ATC AAG CGG CGC TCG CAG TCC arg val leu gly ser val ser gln phe glu glu phe gly arg val AGG GTG CTG GGC AGT GTG TCC CAG TTT GAG GAG TTT GGC CGT CC GTG TGG GGC AGT GTG TCC CAG TTT GAG GAG TTT GGC CCT CC GTG TGG GGC AGT GTG TCC CAG TTT GAG GAG TTT GGC CCT GGT GCC AGG CTG CTG CTG CTG CTG CTG CCC CCA CTG GGT GCC AGG CTG CTG CTG CTG CTG CTT CCA GCC CCT CAG GGT GGG CCC CCC TTG GGC ACC TGC CTT CCA GCC CCT CAG GGT GAG CTT TGG GGG GCT TTT GCT GCA ATA TAC TGG GGA TTG TAC GGG CCC CAC CTT CGC TGT GCT GCT CTT CAG GTT TGC CTC CTC CTT CGC TTT GCT GCT ATA TAC TGG					leu TIA					
919 91h tyr asp arg ser gly asn leu leu his trp trp thr GGC CAG TAT GAC CCC TCA GGG AAC CTG CTG CAC TGG TGG AGG leu tyr asp asn phe thr val tyr asn gln arg val asn gly ccc TAT CAC TAT CAC TAT CAC TAT CAC CAG GTG CAG GTG CAG GCC TAT CAC AAG TGG GTG CGG GAG CAC GGC CAA ile ala phe ala gln asn trp cys ile lys arg arg ser gln arg ccc TTT GCC CAG AAC TGG TGC ATC AAG GGG CGG TGC CAG ATC TAT GCC TTT GCC CAG TTT GCC CCC CCG GGG CGG TTT GCC CAG TTT GAG GAG TTT GCC CCT CCC GCG GGG TTT GGG GGG TTT GCC CCT TCC GTG TGG TG					val					
919 91h tyr asp arg ser gly asn leu leu his trp trp GGC CAG TAT GAC CGC TCA GGG AAC CTG CTG CAC TGG TGG TGG TAT GAC TAT ACT GTC TAC AAC CTG CGG GTG AAC ala tyr dan thr val tyr asn gln arg val asn GCC TAT CAC AAC TGG GTG CGG GAG CAC GGC TAT CAC AAC TGG TGC GTG CGG GAG CAC GGC TAT GCC TAT CAC AAC TGG TGC TAC AGG GGG CGG TGG ATT GCC TTT GCC CAG TAT GAC GGG CGG TGG TGG TAT GCC TTT GCC CAG TTT GCC CCA GGG CGG TTT GCC CAG TTT GAG GAG TTT GCC CAG TTT GAG GAG TTT GCC CCA GGT GTG TGC CTG CCC CCA GGT GTG TGC CTG CCC CCA GGT GTG TGC CTG CCC CCA GGT GTG TGG CTG CTG CTG CTG TGG TGG TTT GGG GAA ATA GGT GTG TAC GGG CCT TGC CTT CCA GCC CAG TTG TAC GGG CTT CTT GCT GTT CTT GCT GCT GCA TTG TAC TGT TAC GGC CCC CAC TTT TGG GGG GCT GTT GTT GTT GTT GTT GCT GC					arg CGG		CTG			
919 91n tyr asp arg ser gly asn leu leu his trp GGC CAG TAT GAC CGC TCA GGG AAC CTG CTG CAC TGG CTG CTG CTG CTG CTG CTG CTG CTG CT					91 <u>y</u> 660		CCA			
919 91h tyr asp arg ser gly ash leu leu his GGC CAG TAI GAC CGC TCA GGG AAC CTG CTG CAC leu tyr ash gln arg CTC TAI GAC AAC TTC ACT GTC TAC AAC CAG CGG ala tyr his ala tyr gln lys trp val arg glu GCC TAC CAC GCC TAI CAG AAG TGG GTG CGG GAG ile ala phe ala gln ash trp cys ile lys arg ATT GCC TTT GCC CAG AAC TGG TGC ATC AAG GGG ATT GCC TTT GCC CAG TCC ACC TTT GCC GGG GAG TCC GTG TGC TTT GAG GAG GTG CTG TGG TGC TTT GAG GAG GTG GTG TGG TGG TGG TGC TGC TGC TGC CTT CC GTG TGG TG					phe		SSS			
919 91n tyr asp arg ser gly asn leu leu GGC CAG TAT GAC CGC TCA GGG AAC CTG CTG LAU tyr asn gln CTC TAT GAC AAC TTC ACT GTC TAC AAC CAG ala tyr gln lys trp val arg GCC TAT CAC AAG TGG GTG CGG Ile ala phe ala gln asn trp cys ile lys ATT GCC TTT GCC CAG AAC TGG TGC ATC AAG ATG GTG CGC CGC TAT GCC TTT GCC CAG TTT GAG Ser val ser gln phe glu AGG GTG GGG AGT GTG TCC CAG TTT GAG GGT GTG TGG TGG TGG TGC TGC GCC TGC GGT GTG GGG GCT GGG GCT GTG GGT GTG GGG GCT GTG GGA TTG TAG GGG GCT GTG GGA TTG TAG GGG GCT GTG GGA TTG TAG GGG CCC CAC TTT GGG GCT GTG GGA TTG TAG GGG CCC CAC CTT CGC TGT GTT	his CAC						ACG	CIT		
919 91h tyr asp arg ser gly ash GGC CAG TAT GAC CGC TCA GGG AAC leu tyr asp asn phe thr val tyr CTC TAT GAC TAC TTC ACT GTC TAC ala tyr gln lys trp GCC TAT GAC TAC TAG AAC TGG TGC ATT GCC TAT GCC CAG AAC TGG TGC ATT GCC TTT GCC CAG AAC TGG TGC AGG GTG GTG GTG GTG GTG GTG GTG GTG GTG	leu CTG	gln CAG		lys Aag	glu GAG		IGC	TGC		
919 91n tyr asp arg ser gly asn GGC CAG TAT GAC CGC TCA GGG AAC leu tyr asp asn phe thr val tyr CTC TAT GAC AAC TTC ACT GTC TAC ALA TYR GCC TAT CAG AAG TGG TGC TAT GCC TAT GCC TAT GCC TAT GCG AAG TGG ATG TGC TAT GCC TAT GCC TAT GCC CAG AAC TGG TGC TGC TGC GGC GGC GGC GGC GGC GG	leu Ciù		val GIG	ile ATC	pha TTT		ည္သ	ACC		
919 91n tyr asp arg ser GGC CAG TAT GAC CGC TCA leu tyr asp asn phe thr CTC TAT GAC AAC TTC ACT ala tyr gln GCC TAC CAC GCC TAT CAG GCC TAT CAG ATT GCC TTT GCC CAG AAC ATT GCC TTT GCC CAG ACT GCG GGC AGT GTG TTC GTG TGG GGC GCT GGG CCC CAC GGG CCC CAC	AAC			0 <u>7</u> 3	gla		ပ္ပ	ACC		ည္သ
919 91n tyr asp arg ser GGC CAG TAT GAC CGC TCA leu tyr asp asn phe thr CTC TAT GAC AAC TTC ACT ala tyr gln GCC TAC CAC GCC TAT CAG GCC TAT CAG ATT GCC TTT GCC CAG AAC ATT GCC TTT GCC CAG ACT GCG GGC AGT GTG TTC GTG TGG GGC GCT GGG CCC CAC GGG CCC CAC	917 966			136 136	ser TCC		CIG	ညည		
919 91n tyr asp arg GGC CAG TAT GAC CGC leu tyr asp asn phe CTC TAT GAC AAC TTC GCC TAC CAC GCC TAT ile ala phe ala gln ATT GCC TTT GCC CAG arg val leu gly ser AGG GTG CTG GGC AGT TTC GTG TGG GGC GCC GGT GCG GGC GCT CCG GGC GGC GCT GGT GGC TGG GCT GGT GGC TGG GGC GGT GTG TGG GCT GGT GGC TAC GGC GGT GAG GCT GGA GGA TTG TAC GGG CCC	age TG			asn	val		1 GG	CIG		CAC
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	asp GAC		ala GCC	ala GCC	91 <u>y</u> 660	ter	TGA	CCC	CCT	999
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	25 25 25 25 25 25 25 25 25 25 25 25 25 2					val	o Fig	ပ္ပင္ပ	GGT	IIG
2071 2161 2341 2431 2521 2611 2701								GGT	CAG	GGA
	2071	2161	2251	2341	2431	,	2521	2611	2701	
										-

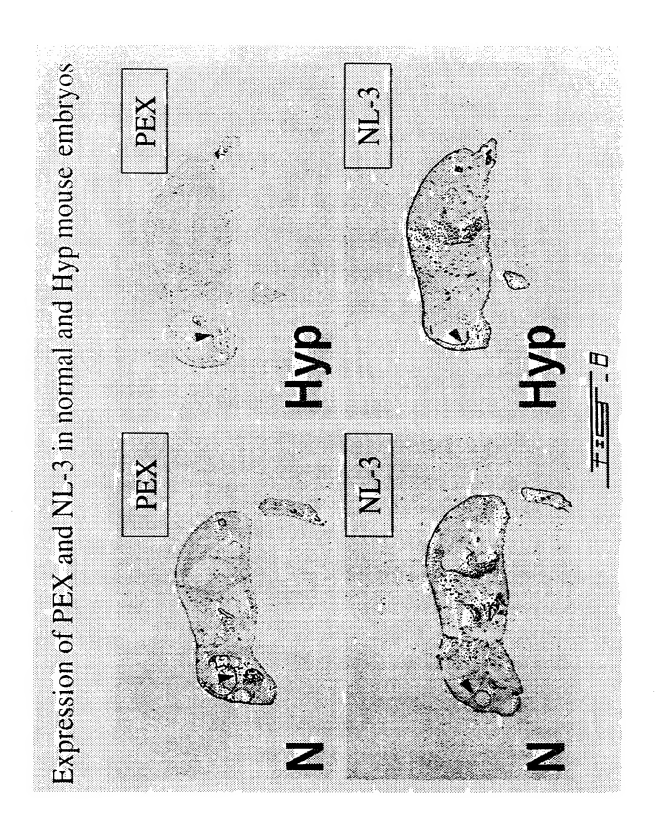
fr = 5 (cont'd)

	Sequence comparison between NEP, NL1, NL2 and NL3
NEP-HUM	1 10 20 30 40 MGKSESQMDITDINTPKPKKKQRWTPLEISLSVLVLLLTIIAV
NL1-MOU	MVERAGWCRKKSPGFVEYGLMVLLLLLLGAIVTLG.VFYSI.GKQL
NL2-HUM	
NL3-HUM	in the second
NEP-HUM	50 60 70 TMIALYATYDDGICKSSDCIKSAARLIQ.NMDATT
NL1-MOU	PLLTSLLHFSWDERTVVKRALRDSSLKSDICTTPSCVIAAARILE.NMDQSR
NL2-HUM	PRLASRLCFLQEERTFVKRKPRGIPEAQEVSEVCTTPGCVIAAARILQ.NMDPTT
NL3-HUM	CLLSGLVFAAGLCAILAAMLALKYLGPVAAGGGACPEGCPERKAFARAARFLAANLDASI
NEP-HUM	80 90 100 110 120 130 EPCTDFFKYACGGWLKRNVIPETSSRYGNFDILRDELEVVLKDVLQEPKTEDIVAVQ.KA
NL1-MOU	NPCENFYQYACGGWLRHHVIPETNSRYSVFDILRDELEVILKGVLEDSTSQHRPAVE.KA
NL2-HUM	EPCDDFYQFACGGWLRRHVIPETNSRYSIFDVLRDELEVILKAVLENSTAKDRPAVE.KA
NL3-HUM	DPCQDFYSFACGGWLRRHAIPDDKLTYGTIAAIGEQNEERLRRLLARPGGGPGGAAQRKV
NEP-HUM	140 150 160 170 180 190 KALYRSCINESAIDSRGGEPLLKLLPDIYGWPVATENWEQKYGASWTAEKAIAQLNSKYG
NL1-MOU	KTLYRSCMNQSVIEKRDSEPLLSVLKMVGGWPVAMDKWNETMGLKWELERQLAVLNSQFN
NL2-HUM	RTLYRSCMNQSVIEKRGSQPLLDILEVVGGWPVAMDRWNETVGLEWELERQLALMNSQFN
NL3-HUM	RAFFRSCLDMREIERLGPRPMLEVIEDCGGWDLGGAEERPGVAARWDLNRLLYKAQGVYS
NEP-HUM	200 210 220 230 240 250 KKVLINLFVGTDDKNSVNHVIHIDQPRLGLPSRDYYECTGIYKEACTAYVDFMISVARLI
NL1-MOU	RRVLIDLFIWNDDQNSSRHVIYIDQPTLGMPSREYYFQEDNNHKVRKAYLEFMTSVATML
NL2-HUM	RRVLIDLFIWNDDQNSSRHIIYIDQPTLGMPSREYYFNGGSNRKVREAYLQFMVSVATLL
NL3-HUM	AAALFSLTVSLDDRNSSRYVIRIDQDGLTLPERTLYLAQDEDSEKVLAAYRVFMERVL

	260 270 280 290 300 310
NEP-HUM	200 200 310
NL1-MOU	RKDQNLSKESAMVREEMAEVLELETHLANATVPQEKRHDVTALYHRMDLMELQERFGL
NL2-HUM	REDANLPRDSCLVQEDMVQVLELETQLAKATVPQEERHDVIALYHRMGLEELQSQFGL
NL3-HUM	SLLGADAVEQKAQEILQVEQQLANITVSEYDDLRRDVSSMYNKVTLGQLQKITP.
NEP-HUM	320 330 340 350 360 370 EINGKPFSWLNFTNEIMSTVNISITNEEDVVVYAPEYLTKLKPILTKYSARDLQNLMSWR
NL1-MOU	KGFNWTLFIQNVLSSVEVELFPDEEVVVYGIPYLENLEDIIDSYSARTMQNYLVWR
NL2-HUM	KGFNWTLFIQTVLSSVKIKLLPDEEVVVYGIPYLQNLENIIDTYSARTIQNYLVWR
NL3-HUM	HLRWKWLLDQIFQEDFSEEEEVVLLATDYMQQVSQLIRSTPHRVLHNYLVWR
NEP-HUM	380 390 400 410 420 430 FIMDLVSSLSRTYKESRNAFRKALYGTTSETATWRRCANYVNGNMENAVGRLYVEAAFAG
NL1-MOU	* *** ** * ** ****** * * * * * * * * *
NL2-HUM	LVLDRIGSLSQRFKDTRVNYRKALFGTMVEEVRWRECVGYVNSNMENAVGSLYVREAFPG
NL3-HUM	* * * * * * * * * * * * * * * * * * *
NEP-HUM	440 450 460 470 480 490 ESKHVVEDLIAQIREVFIQTLDDLTWMDAETKKRAEEKALAIKERIGYPDDIVSNDNK.L
NL1-MOU	DSKSTVRELIEKIRSVFVDNLDELNWMDEESKKKAQEKAMNIREQIGYPDYILEDNNKHL
NL2-HUM	DSKSMVRELIDKVRTVFVETLDELGWMDEESKKKAQEKAMSIREQIGHPDYILEEMNRRL
NL3-HUM	ASKAKVQQLVEDIKYILGQRLEELDWMDAETRAAARAKLQYMMVMVGYPDFLLKPDAV
NEP-HUM	500 510 520 530 540 550 NNEYLELNYKEDEYFENIIQNLKFSQSKQLKKLREKVDKDEWISGAAVVNAFYSSGRNQI
NL1-MOU	DEEYSSLTFYEDLYFENGLQNLKNNAQRSLKKLREKVDQNLWIIGAAVVNAFYSPNRNOI
NL2-HUM	DEEYSNLNFSEDLYFENSLQNLKVGAQRSLRKLREKVDPNLWIIGAAVVNAFYSPNRNQI
NL3-HUM	DKE.YEFEVHEKTYFKNILNSIRFSIQLSVKKIRQEVDKSTWLLPPQALNAYYLPNKNQM
	TE [(cont'd)

NEP-HUM	560 570 580 590 600 610 VFPAGILQPPFFSAQQSNSLNYGGIGMVIGHEITHGFDDNGRNFNKDGDLVDWWTQQS.	AS
NL1-MOU	VFPAGILQPPFFSKDQPQSLNFGGIGMVIGHEITHGFDDNGRNFDKNGNMLDWWSNFS	AR
NL2-HUM	VFPAGILQPPFFSKEQPQALNFGGIGMVIGHEITHGFDDNGRNFDKNGNMMDWWSNFS	TQ
NL3-HUM	VFPAGILQPTLYDPDFPQSLNYGGIGTIIGHELTHGYDDWGGQYDRSGNLLHWWTEAS	YS
NEP-HUM	620 630 640 650 660 670 NFKEQSQCMVYQYGNFSWDLAGGQHLNGINTLGENIADNGGLGQAYRAYQNYIKKNGER	EK
NL1-MOU	HFQQQSQCMIYQYGNFSWELADNQNVNGFSTLGENIADNGGVRQAYKAYLRWLADGGKI	+ +
NL2-HUM	HFREQSECMIYQYGNYSWDLADEQNVNGFNTLGENIADNGGVRQAYKAYLKWMAEGGKI	DQ
NL3-HUM	RFLRKAECIVRLYDNFTVYNQRVNGKHTLGENIADMGGLKLAYHAYQKWVREHGPE	ЕН
NEP-HUM	680 690 700 710 720 730 LLPGLDLNHKQLFFLNFAQVWCGTYRPEYAVNSIKTDVHSPGNFRIIGTLQNSAEFSEA	۱F
NL1-MOU	RLPGLNLTYAQLFFINYAQVWCGSYRPEFAVQSIKTDVHSPLKYRVLGSLQNLPGFSEA	\F
NL2-HUM	QLPGLDLTHEQLFFINYAQVWCGSYRPEFAIQSIKTDVHSPLKYRVLGSLQNLAAFADT	'F
NL3-HUM	PLPRLKYTHDQLFFIAFAQNWCIKRRSQSIYLQVLTDKHAPEHYRVLGSVSQFEEFGRV	'L
NEP-HUM	740 750 HCRKNSYMNPEKKCRVW	
NL1-MOU	HCPRGSPMHPMKRCRIW	
NL2-HUM	HCARGTPMHPKERCRVW	
NL3-HUM	HCPKVSPMNPAHKCSVW	



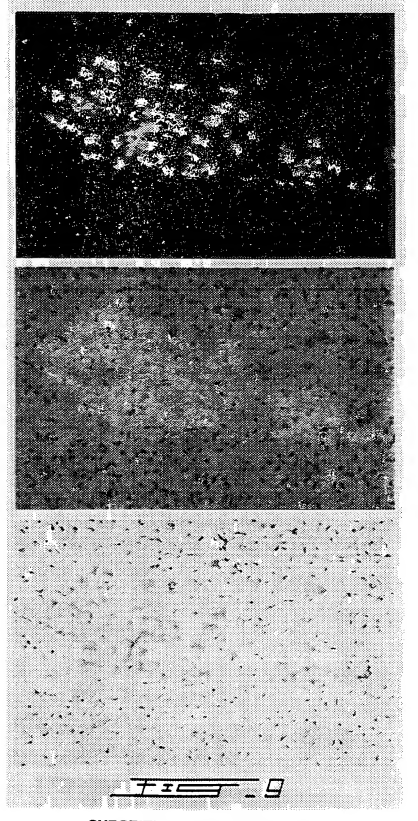


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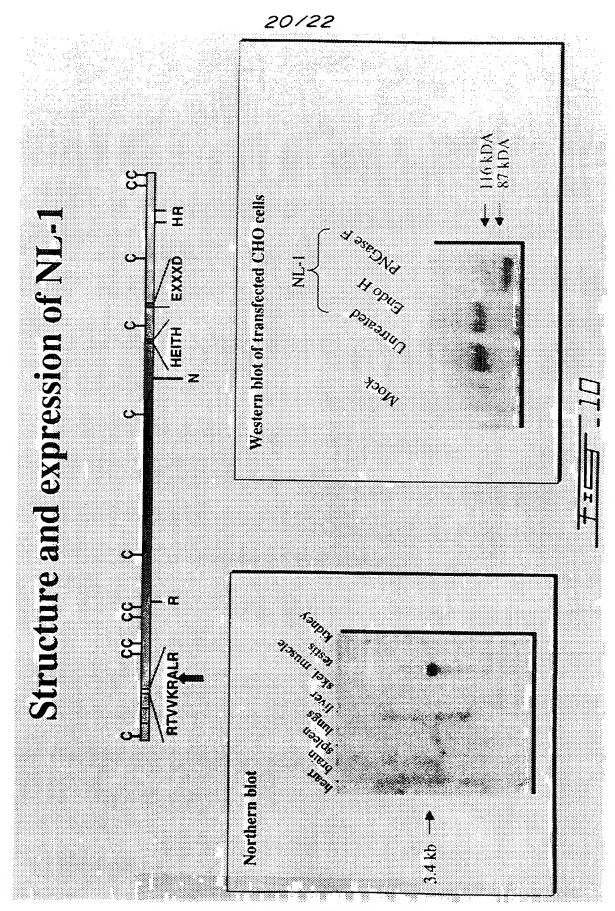
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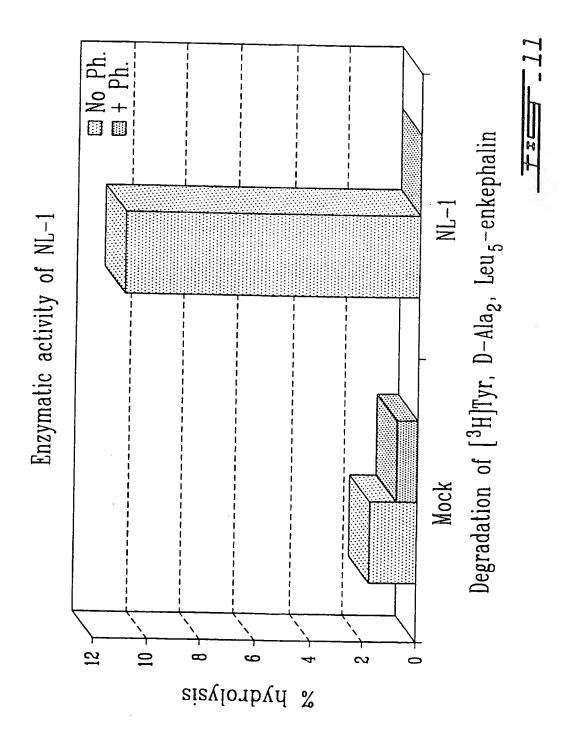
NL3 in the BRAIN

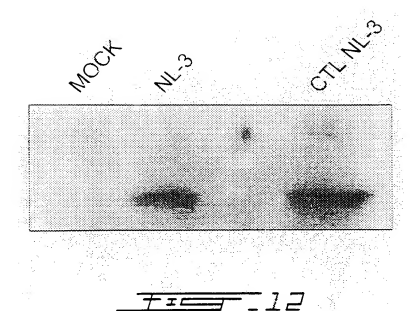


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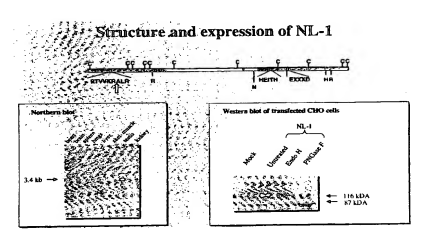
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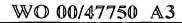
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[Continued on next page]

(54) Title: METALLOPROTEASES OF THE NEPRILYSIN FAMILY



(57) Abstract: In this paper, we describe RT-PCR strategies that allowed us to identify and clone members of the NEP-like family. Degenerate oligoncleotide primers corresponding to consensus sequences located on either side of the HEXXH consensus sequence for zincins were designed and used in RT-PCR with mouse and human testis cDNAs. DNA fragments with lengths expected from the sequence of this class of enzympes were obtained. These DNA fragments were cloned and sequenced. Using this PCR strategy and the PCR fragments as probes to screen cDNA libraries, three zincin-like peptidases were identified in addition of known members of the family. The cDNA sequences allowed to derive specific probes for Northern and in situ hybridization, and probe human chromosomes to localize the gene and establish potential links to genetic diseases. Furthermore, these cDNA sequences were used to produce recombinant fusion proteins in Escherichia coli in order to raise specific antibodies. Finally, the cDNA sequences were cloned in mammalian expression vectors and transfected in various mammalian cell lines to produce active recombinant enzymes suitable for testing specific inhibitors.





Date of publication of the amended claims:

25 May 2001

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

AMENDED CLAIMS

[received by the International Bureau on 28 November 2000 (28.11.00); new claims 23-27 added; remaining claims unchanged (4 pages)]

- 1. A neutral endopeptidase-like metallopeptidase named NL-1 which is isolated by probing tissue nucleic acids with degenerate oligonucleotides derived from a conserved sequence located on either side of a sequence His-Glu-Xaa-Xaa-His, wherein Xaa is any amino acid, which has a sequence selected from the amino acid sequences shown in Figures 3, 4, and 5, and a variant thereof sharing about 80% homology with said sequence.
- 2. A metallopeptidase as defined in claim 1, which has the amino acid sequence shown in Figure 3.
- 3. A metallopeptidase as defined in claim 1, which has the amino acid sequence shown in Figure 4.
- 4. A metallopeptidase as defined in claim 1, which has the amino acid sequence shown in Figure 5.
- 5. A nucleic acid encoding the metallopeptidase of any one of claims 1 to 3.
- A recombinant vector comprising the nucleic acid defined in claim 4.
- 7. A recombinant host expressing the nucleic acid of claim 4.
- 8. A method for producing a metallopeptidase as defined in any one of claims 1 to 3, which comprises the step of culturing a recombinant host as defined in claim 6 in growth supportive medium and recovering said metallopeptidase from the cell or the culture medium.
- A method for screening new molecules related to neural endopeptides, which comprises the step of:
 - aligning nucleotidic sequences of known molecules related to NEP;
 - assessing consensus sequences on either side of a sequence comprising His-Glu-Xaa-Xaa-His sequence;
 - synthetising degenerate sequences of said consensus sequences;

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- contacting said degenerate sequences with the nucleic acids of panel of candidate samples susceptible to express said new molecules, in conditions such that a hybridization complex can form between the nucleic acids of samples and the degenerate sequences;
- detecting said hybridization complex as an indication of a candidate sample which comprises a molecules related to NEP;
- sequencing the nucleic acids of said hybridized complex;
 whereby a new sequence sharing homology with NEP is a new molecule relate to NEP.
- 10. A method as defined in claim 8 wherein said degenerate sequences are selected from Figure 2.
- 11. A method for producing a soluble form of a membrane protein of interest having a C-terminal ectodomain, said soluble form essentially consisting a said ectodomain, which comprises:
 - obtaining nucleic acids encoding essentially the ectodomain;
 - fusing the nucleic acids in phase with an amino terminal fragment of NL-1 or NL-2 as defined in C;
 - having the fused nucleic acids to be expressed in a host cell in the presence of a culture medium, which expresses or is made to express furin;
 - recovering said soluble form in the culture medium.
- 12. A method as defined in claim 9, wherein said protein of interest is NL-3 or β -endorphin.
- 13. An oligonucleotide selected from those in Figure 2.
- 14. A composition of matter comprising one or more of those in Figure 2.
- 15. A recombinant vector comprising a nucleic acid encoding the N-terminal part of the amino acid sequence shown in Figure 3 or 4 which N-terminal part terminates with a furin-recognition sequence.
- 16. A host transformed with the recombinant vector of claim 13.

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- 17. The soluble form of the metallopeptidase defined in any one of claims 1 to 3 which soluble form essentially consist of the ectodomain of said metallopeptidase.
- 18. A composition comprising the soluble metallopeptidase of claim 15.
- An oligonucleotide derived from the nucleic acid defined in claim 4, which oligonucleotide has at least 12 nucleic acid in length.
- 20. An antibody directed against the metallopeptidase defined in any one of claims 1 to 3.
- 21. A method for detecting the presence of the metallopeptidase as defined in any one of claims 1 to 3 in a sample, which comprises the steps of contacting: contacting said sample with the antibody defined in claim 18, in conditions such that an immune complex is formed between said antibody and said metallopeptidase, and detecting the presence of an immune complex as an indication of the presence of said metallopeptidase in said sample.
- 22. A method of detecting the presence or amount of the metallopeptidase as defined any one of claims 1 to 3 in a sample, which comprises the steps of: contacting said sample with the nucleic acid defined in claim 4 or with an oligonucleotide as defined in claim 17 in conditions such that a hybridization complex is formed between the target nucleic acids of the sample and the nucleic acids or oligonucleotides encoding said metallopeptidase, and detecting the formation of such hybridization complex as an indication of the presence of said metallopeptidase in said sample.
- 23. A method for obtaining an inhibitor of a neutral endopeptidase-like enzyme (NEP-like), which comprises the steps of :
 - -- contacting NEP-like with a molecule or extract in the presence of a NEP-like substrate; and
 - assaying the resulting solution for the intact substrate or for a decrease in the hydrolysed substrate as an indication of the presence of said inhibitor.

- 24. A method as described in claim 23, wherein said NEP-like has an amino acid sequence chosen from the amino acid sequences shown in Figures 3, 4 and 5 or a variant or fragment thereof.
- 25. A method as described in claim 23 or 24, wherein said NEP-like substrate is Tyrosyl-[3,5-3H])(D-Ala₂)-Leu₅-enkephalin or bradykinin.
- 26. A method as described in any one of claims 23 to 25, wherein said assaying is realised with specific antibodies, HPLC or by the appearance of fluorescence when a self-quenched fluorescence tagged peptide is used as said NEP-like substrate.
- 27. A method as defined in any one of claims 23 to 26, wherein said molecule or extract is selected from identified synthetic libraries, biota extracts and from rationally designed inhibitors using X-ray crystallography and substituent activity relationships.

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